

ERDEC-SP-054

OF NEUTRALIZED CHEMICAL AGENT IDENTIFICATION SETS

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NON-STOCKPILE CHEMICAL MATERIEL PROGRAM

August 1997

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19971209 059

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REPORT DOCUMENTATION PAGE

Form Approved
OMB No. 0704-0188

Public reporting burden for this collection of information is estimated to average 1 hour per response, including the time for reviewing instructions, searching existing data sources, gathering and maintaining the data needed, and completing and reviewing the collection of information. Send comments regarding this burden estimate or any other aspect of this collection of information, including suggestions for reducing this burden, to Washington Headquarters Services, Directorate for Information Operations and Reports, 1215 Jefferson Davis Highway, Suite 1204, Arlington, VA 22202-4302, and to the Office of Management and Budget, Paperwork Reduction Project (0704-0138), Washington, DC 20503.

Davis Highway, Suite 1254, Armington, VA 22222 1555	T	T	DATE COVERED
1. AGENCY USE ONLY (Leave blank)	2. REPORT DATE	3. REPORT TYPE AN	
	1997 August	Final, 95 Nov	
4. TITLE AND SUBTITLE			5. FUNDING NUMBERS
Evaluation of the Vesicat		eutralized	None
Chemical Agent Identifica	tion Sets		
6. AUTHOR(S)			
Olajos, Eugene J.; Salem	Harry (ERDEC); and (Gieseking,	
John K. (PMNSCM)			·
7. PERFORMING ORGANIZATION NAME	(S) AND ADDRESS(ES)		8. PERFORMING ORGANIZATION
DIR, ERDEC, ATTN: SCBRD-R		23	REPORT NUMBER
PM, Non-Stockpile Chemica			ERDEC-SP-054
ATTN: SFAE-CD-N/NP/NM, A	APG. MD 21010-5401		Elded by 034
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9. SPONSORING/MONITORING AGENCY	ANAME(S) AND ADDRESS(ES)		10. SPONSORING / MONITORING
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11. SUPPLEMENTARY NOTES			
12a. DISTRIBUTION / AVAILABILITY STA	TEMENT		12b. DISTRIBUTION CODE
Approved for public relea	se; distribution is	unlimited.	1
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13. ABSTRACT (Maximum 200 words)			
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14. SUBJECT TERMS Acute toxicity (dermal) Microvesication Microblister	Guinea-pig (hairless) Wastestreams Rapid Response System Agent (HD,HN, L)	Sulfur mustard CAI Nitrogen mustard Lewisite Chemical neutralization	15. NUMBER OF PA	AGES
Histopathology 17. SECURITY CLASSIFICATION OF REPORT UNCLASSIFIED	RRS 18. SECURITY CLASSIF OF THIS PAGE UNCLASSIFIED	DCDMH FICATION 19. SECURITY CLASSIFI OF ABSTRACT UNCLASSIFIED	CATION 20. LIMITATION OF	ABSTRACT

(DCDMH)].

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Study Number: G155538A

This Study was inspected by the Quality Assurance Unit and reports were submitted to the Study Director and management as follows:

Phase Inspected	Inspection Date	Dated Reported to Study Director	Date of Report to Management
Dilution	2/14/96	3/4/96	3/4/96
CSM decontamination	2/19/96	3/4/96	3/4/96
Test article administration - dermal	2/19/96	3/4/96	3/4/96
Test system preparation	2/19/96	3/4/96	3/4/96
Gas chromotography analysis	2/20/96	3/4/96	3/4/96 .
Sample collection	2/20/96	3/4/96	3/4/96
Histology processing	2/22/96	3/4/96	3/4/96
Test system preparation	6/26/96	7/1/96	7/1/96
Test article administration - dermal	6/26/96	7/1/96	7/1/96
Histology processing	6/27/96	7/1/96	7/1/96
Necropsy/tissue collection	6/27/96	7/1/96	7/1/96
Euthanasia	6/27/96	7/1/96	7/1/96
Histology processing	6/28/96	7/1/96	7/1/96
Audit study file	8/9/96	8/9/96	9/16/96
Audit study file	10/8/96	10/8/96	12/5/96
Audit study file	10/15/96	10/15/96	11/14/96
Audit study file	1/7/97	1/7/97	2/28/97
Audit study file	4/4/97	4/4/97	4/25/97
Protocol	5/16/97	5/16/97	5/29/97
Draft Final Report	5/16/97	5/16/97	5/29/97
Audit study file	5/27/97	5/27/97	5/29/97
Final Report	6/5/97	6/5/97 Eliston MREF Quality	Assurance Unit, Date

GLP COMPLIANCE STATEMENT

The percutaneous dosing of hairless guinea pigs with wastestreams, neutralizing solution and known vesicants, and the gross and histopathologic evaluations of skin lesions in this study were performed by Battelle in compliance with the Environmental Protection Agency's (EPA) Good Laboratory Practice (GLP) Standards (40 CFR Part 792). Likewise, evaluation of the analytical method for HD, HN-1 and L in wastestreams and the determination of HD or HD, HN-1 and L concentrations, as appropriate, in wastestreams was accomplished at Battelle in compliance with EPA GLP Standards. Reports on findings from searches of the literature on HD, HN-1 and L degradation and degradation products and their vesicancy potential as well as analyses of wastestreams for degradation products and residual agent concentrations performed elsewhere than the MREF are excepted from this Good Laboratory Practices Compliance Statement. This study was conducted according to the study protocol, as amended, and Battelle's standard operating procedures. Deviations from the protocol or standard operating procedures are documented in Appendix A. The data presented accurately reflect the results of this study.

Carl T. Olson, D.V.M., Ph.D.

Study Director

Date

QUALITY ASSURANCE

The analytical data supplied by the U. S. Army Edgewood Research, Development and Engineering Center (ERDEC) in support of this task were generated under the auspices of the Research and Technology Directorate Quality Assurance Program Plan. Accordingly, the data are supported by written methodology, sample identification records, and suitable instrument maintenance and calibration. The data and supporting records are retained by ERDEC.

DENNIS W. JOHNSON

Quality Assurance Coordinator

Research and Technology Directorate

REPORT REVIEW

The report entitled, "Vesication Evaluation of Neutralized Chemical Agent Identification Sets (CAIS)" was reviewed for technical accuracy in data analysis and report approach. To the best of our knowledge, the report was considered to be an accurate reflection of the vesication data presented in the original report by Olson et al. (1997) titled "Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Set (CAIS) Components".

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LIST OF ACRONYMS AND ABBREVIATIONS

CAIS Chemical Agent Identification Sets

CASARM Chemical Agent Standard Analytical Reference Materiel

CFR Code of Federal Regulations

CI Chemical Ionization
CW Chemical Warfare

DCDMH 1, 3-Dichloro-5,5 Dimethylhydantoin

DoD Department of Defense

DOT Department of Transportation

El Electron Impact

EPA Environmental Protection Agency

ERDEC U.S. Army Edgewood Research, Development and Engineering Center

GC/MS Gas Chromotagraphy/Mass Spectrometry

GLP Good Laboratory Practice

H Sulfur Mustard HD Sulfur Mustard

HMR Hazardous Materials Regulations

HMTA Hazardous Materials Transportation Act

HN Nitrogen Mustard HS Sulfur Mustard LD₁₀ Lethal Dose Low

L Lewisite

LC₅₀ Lethal Concentration 50

LD₅₀ Lethal Dose 50

NSCM Non-Stockpile Chemical Materiel

NSCMP Non-Stockpile Chemical Materiel Program

PMCD Program Manager for Chemical Demilitarization
PMCSD Project Manager for Chemical Stockpile Disposal
PMNSCM Project Manager for Non-Stockpile Chemical Materiel

ppm Parts per Million

RRS Rapid Response System t-BuOH Tertiary-butyl Alcohol

TD_{to} Toxic Dose Low

TSDF Treatment, Storage, and Disposal Facility

PREFACE

The work described in this report was authorized under the Chemical Demilitarization Program. This work was started in November 1995 and completed in August 1997.

The use of either trade or manufacturers' names in this report does not constitute an official endorsement of any commercial products. This report may not be cited for purposes of advertisement.

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Evaluation of the Vesicating Properties of Neutralized Chemical Agent Identification Sets

1. INTRODUCTION

The U.S. Army Program Manager for Chemical Demilitarization (PMCD) has been designated as the single agency within the Department of Defense (DoD) to destroy all chemical warfare-related materiel. Destruction of chemical weapons which are part of the unitary stockpile is the responsibility of the Project Manager for Chemical Stockpile Disposal (PMCSD). The demilitarization/destruction of non-stockpile chemical materiel (NSCM), among those items are Chemical Agent Identification Sets (CAIS), is the responsibility of the Project Manager for Non-Stockpile Chemical Materiel (PMNSCM). CAIS may contain agent in chloroform (HD, HN or L), agent (HD, HN or L) adsorbed on charcoal, and agent (HD) in neat form - all packed in glass ampules. CAIS may also contain miscellaneous materiel/industrial chemicals. CAIS were declared obsolete in 1971.

The PMNSCM is developing the Rapid Response System (RRS) for processing CAIS. The RRS is a system of trailer-mounted equipment designed to support on-site characterization and primary treatment of recovered CAIS. The system is designed for unpacking, identification of chemicals, segregation of CAIS components, neutralization of the chemical agents, repackaging of industrial chemicals. The chemical agent wastes and repacked industrial chemicals will be provided to a hazardous waste treatment, storage, and disposal facility (TSDF) for ultimate disposal.

The RRS is designed to provide a safe and environmentally secure work area to chemically treat mustard and lewisite agents, and to repackage industrial chemicals. The containers of neutralized mustards and lewisite, dunnage, neutralents, and the repacked industrial compounds will be transferred to a TSDF for disposal. It should be noted that the primary objective of the RRS chemical neutralization process is to convert CAIS chemical agents to less toxic products to minimize the health hazard associated with the handling and transportation of demilitarized non-stockpile chemical materiel. Given the chemistry and toxicity characteristics of the chemical agents/degradation products, and the reactants used in the neutralization process, the wastestreams designated for transportation to TSDFs are expected to be complex mixtures exhibiting various degrees of acute dermal toxicities such as irritation; however, other skininjurious effects (i.e. vesication) are possible.

In preparation for the RRS test to be conducted at Deseret Chemical Depot, Utah, the PMNSCM has conducted dermal toxicity evaluations (vesication testing) of wastestreams resultant from the chemical neutralization of CAIS to determine reduction of agent and/or agent degradation product vesicancy. This document is a synopsis of a technical report (Olson et al, 1997) on results of vesicancy studies conducted in hairless guinea-pigs.

1.1 PROCESS OVERVIEW

Process chemistry developed for chemical agent "detoxification", referred to as the neutralization process, focused on chemical methods that were capable of converting chemical agents to products/by-products with marked reduction in agent characteristics (i.e. vesication). Thus, chemical neutralization processes were sought which: (1) achieved process simplicity, (2) resulted in marked reduction in agent characteristics, and (3) generated wastestreams having reduced toxicity characteristics that can be handled and disposed of in a manner similar to industrial chemicals and/or wastes. The process combines chemical agent(s) with a treatment solution, and oxidizing chemical dissolved in an organic/aqueous solvent mixture. The final product of the chemical neutralization process (wastestream) is a complex mixture composed of reaction products, by-products, unreacted excess reactants, and residual chemical agent(s).

A moderate oxidizing agent, 1, 3-dichloro-5, 5-dimethylhydantoin (DCDMH), was used in all RRS process chemistries. The reaction conditions varied depending on stoichiometry, sequence of addition of reacting mixtures, and physical condition of the chemical agent - whether neat, dissolved in chloroform, or adsorbed on charcoal.

In the nomenclature of the RRS, chemical treatment of neat HD with DCDMH in organic/aqueous solvent is referred to as the modified "Blue" process. Chemical neutralization of dilute chloroform solutions containing HD, HN-1 or L with DCDMH in organic/aqueous solvent is identified as the modified "Red" process. Chemical neutralization of HD, HN-1 or L adsorbed on charcoal with DCDMH in organic/aqueous solvent is referred to as the "Charcoal" process. Toxicity characteristics of wastestreams resulting from these neutralization reactions were expected to differ from those of chemical agents and/or severe irritant oxidant/solvent systems used in these reactions. Consequently, the changed toxicity characteristics of the resultant wastestreams are primarily attributed to reduction in the concentration of chemical agent(s). However, the vesicancy potential of the wastestreams would depend on the degradation product(s) profile - presence of vesicating moieties such as HD sulfone and divinyl sulfone.

1.2 STUDY PLAN

Evaluation of the vesicancy potential of neutralized CAIS was conducted using a validated animal model (hairless guinea-pig) to assess vesication. Animals were dosed with test article, and treated skin evaluated for microblister formation using light microscopy. The objective was to determine the efficacy of the neutralization process in reducing the vesicating properties of agent (HD, HN or L) as well as forming product solutions with minimal vesicant potential.

2. Materials and Methods

2.1 Chemicals

2.1.1 Agents

Sulfur mustard [2,2'-dichlorodiethyl sulfide (HD), CAS #505-60-2] furnished from Medical Research and Evaluation Facility (MREF) stocks was used neat (undiluted) as a positive control article for vesication¹. Lewisite [dichloro-2-chlorovinyl arsine (L)] CAS #541-25-3 was also furnished from MREF stock. U.S. Army Edgewood Research, Development and Engineering Center (ERDEC) provided a 20 percent solution of nitrogen mustard [bis (2-chloroethyl) ethylamine (HN-1), CAS #538-07-8] in chloroform. Ten percent solutions of HD, HN-1, or L in chloroform (Pretreated - CAIS which also served as control articles in the Phase II portion (dose-ranging) of the vesication studies) were prepared at the MREF laboratory and evaluated for vesicant activity and other histopathology following dermal application.

2.1.2 Chemical Agent Identification Sets (Synthesized)

Actual ampules from CAIS kits were not used; however "CAIS components" were prepared from agent stocks to contain 10 percent agent in chloroform (Chatfield, et al. 1995). Chemical Agent Standard Analytical Reference Material (CASARM) grade HD CAS# 505-60-2 (97.5 mole %), nitrogen mustard [bis (2-chloroethyl) ethylamine (HN-1)] CAS #538-07-8 (≥97% by weight), and CASARM grade lewisite [dichloro-2-chlorovinyl arsine (L)] CAS #541-25-3 (97.8 % by weight) from stocks maintained by the Operations Directorate, ERDEC were used in the preparation of synthesized CAIS. CASARM for HN-1 is not available.

2.1.3 Neutralized Chemical Agent Identification Sets (Wastestreams)

Wastestreams were provided by ERDEC, Aberdeen Proving Ground, MD. Wastestreams from the chemical neutralization of "CAIS components" prepared from agent stocks were tested for vesicancy potential. These wastestreams were prepared by ERDEC as follows:

¹The chemical agents found in CAIS include sulfur mustard, nitrogen mustard, or lewisite. Sulfur mustard was used as representative vesicant for these blistering agents.

- Wastestreams from the neutralization of neat HD with 1,3-dichloro-5,5-dimethylhydantoin (DCDMH) in CHCl₃/t-BuOH/3% H₂O ("Blue" process).
- Wastestreams from the neutralization of 10% HD, HN, or L (agent in CHCl₃) with DCDMH in CHCl₃/t-BuOH/3% H₂O ("Red" process).
- Wastestreams from neutralization of HD, HN, or L (agent on charcoal) with DCDMH in CHCl₃ (HD, HN samples) and with DCDMH in CHCl₃/t-BuOH/3% H₂O (L sample) ("Charcoal" process).

Two wastestreams ("archived" and "fresh") were prepared for each process - "Blue", "Red", and "Charcoal" - and samples sent to the MREF for analysis of agent content and for vesicancy testing. The stability of the wastestreams under conditions of administration were not determined by MREF personnel. Test articles were "archived" and "fresh" "Blue", "Red", and "Charcoal" wastestreams.

2.1.4 Neutralization Solution

Neutralizing solution was prepared at the MREF to determine the effect on the skin of dosing this solution alone. For testing vesicating potential, a 0.555M 1,3-dichloro-5,5-dimethylhydantoin (FW 197.02) control article neutralizing solution was prepared by adding 10.9g DCDMH to a 50:50 tertiary butanol:chloroform with 3 percent water solution in a 100-mL volumetric flask and adding sufficient volume of the butanol/chloroform/water solution to bring the volume to the 100-mL mark. DCDMH (CAS #118-52-5) was purchased from Aldrich Chemical Company (St. Louis, MO). Chloroform (CAS #67-66-3; GC/Spectro grade) was purchased from Burdick and Jackson (Muskegon, MI), and tertiary-butyl alcohol (CAS #75-65-0; ACS Reagent grade) from J.T. Baker (Phillipsburg, NJ). Distilled water was further purified using a Millipore (Bedford, MA) reverse osmosis system.

2.2 Chemical Neutralization of CAIS

RRS chemical neutralization technologies were developed for neutralization of chemical agents HD, HN and L. The primary objective was to develop processes that convert chemical agents to products/by-products that do not exhibit the highly toxic properties of the agents. Additionally, it was desirous to also reduce agent characteristics (i.e. vesication) of the final product solution (wastestream). However, reduction in vesication is not considered a

²"Archived" "Blue" and "Red" wastestreams were initially analyzed at ERDEC (Oct 95) and re-analyzed for agent residual at the MREF and tested for vesicancy (March 96; August 96). "Charcoal" wastestream initially analyzed at ERDEC (Nov 95) was re-analyzed and test for vesicancy at the MREF (March 96; August 96).

[&]quot;Fresh" wastestreams were prepared and initially analyzed at ERDEC (June 96) and reanalyzed and tested for vesicancy at the MREF (June 96; August 96).

requirement per the Hazardous Materials Regulations (HMR). All process chemistries used 1, 3-dichloro-5, 5-dimethylhydantoin (DCDMH) as neutralizing reagent. The solvent system used in the process chemistries was CHC1₃/t-BuOH/3% H₂0). Formulations of treatment reagent/solvent systems for the chemical neutralization of CAIS are presented in Table 1. The principal differences in the chemical composition of wastestreams, originating from the RRS process chemistry, are primarily due to the physichochemical characteristics of the reactants in the reaction mixtures and the agent undergoing chemical treatment. In the presence of oxidizing agent (DCDMH), the chemical agent(s) undergo oxidation, chlorination, substitution, and/or elimination reactions to yield a mixture of products/by-products. Depending on volume, composition, and reaction conditions -- residual chemical agent, products/by-products, and varying amounts of unreacted excess DCDMH may also be present in the wastestreams (Olajos et al, 1996).

TABLE 1. OXIDIZER/SOLVENT SYSTEM STOICHIOMETRY UTILIZED IN THE MODIFIED "BLUE", "RED", AND "CHARCOAL" PROCESS CHEMISTRIES

- 1 volume of neat HD treated with 20 volumes of 0.555M 1,3-dichloro-5,5-dimethylhydantoin (DCDMH) in CHCl,/t-butanol (50/50) with 3% water by volume ("Blue" Process).
- 1 volume of each 10% HD in CHCl₃, 10% HN in CHCl₃, and 10% L in CHCl₃ treated with 4 volumes of 0.555M 1,3-dichloro-5,5-dimethylhydantoin (DCDMH) in 50/50 CHCl₃/t-butanol with 3% water by volume ("Red" Process).
- 45% by weight HD and HN-1 or charcoal treated with excess 1,3-dichloro-5,5-dimethylhydantoin in CHCl₃ combined with 43% by weight L with excess 1,3-dichloro-5,5-dimethylhydantoin in CHCl₃/t-butanol (50/50) with 3% water by volume ("Charcoal" Process).

2.3 Analytical Methodologies

2.3.1 GC-MS Spectroscopy

Chemically-treated (neutralized) CAIS were analyzed for agent residue levels using full scanning gc-ms spectroscopy. GC-MS spectroscopy was conducted at ERDEC on all wastestreams provided to the MREF, and confirmatory gc-ms analysis was also performed at the MREF prior to conducting the bioassays.

Instrumentation used in the ERDEC analysis of "archived" wastestreams (non-quenched samples) was a Hewlett-Packard 5989B MS engine with Chemstation Data System. Analyses

conducted at both ERDEC and the MREF on "fresh" wastestreams, using quenching and derivatization techniques, utilized a Hewlett-Packard Model 5970B Mass Selective Detector (MSD) with an HP 5890A GC and HP 61034 CMS. For procedural details, the reader is referred to Lucas (1997), Lucas (1996), as provided in the report by Olson et al 1997, and Rosso (1995), as provided in ERDEC-TR-372 (Olajos, et al. 1996). Quantitation was based on internal standardization (internal standard = 1,2,4, 5-tetrachlorobenzene). Calibration standards were as follows: HD (purity 97.%%), HN-1 (purity (96.5%), and L (purity 97.8%).

Product identification of the CAIS wastestreams (archived) was accomplished using GC/MS spectroscopy (EI and CI modes). These studies were performed at ERDEC per procedures described by Rosso and co-workers (Rosso et al. 1995) and documented in the report by Olajos et al. (Olajos et al. 1996).

2.3.2 NMR Spectroscopy

Nuclear magnetic resonance (nmr) spectroscopy analyses of "fresh" wastestreams were conducted at ERDEC as an adjunct to gc-ms analyses. These analyses were performed using a Varian Fourier Transform (FT) nmr spectrometer operated at 200 MHZ for ¹H observation and at 50 MHZ for ¹³C observation. Quantitative data were obtained by digital integration of peak areas.

2.4 Vesicancy Testing

2.4.1 Experimental Design

Studies were conducted which utilized a validated animal model to assess agent-induced vesication of skin (Marlow et al, 1990 and Mershon et al, 1990). Microblister formation in the hairless guinea-pig is analogous to the changes seen in humans (Papirmeister et al, 1984). The degree of vesication was assessed before and after neutralization of agents.

Thirty-five male, hairless guinea-pigs were used in a multiphase study (Phase I analytical; Phase II dose-range; Phase III vesicant assessment of wastestreams) to ascertain the vesicant potential of sulfur mustard (HD), agent/chloroform solutions, and product solutions (wastestreams) from chemically-neutralized CAIS. A synopsis of the experimental design is given in Table 2.

Phase II. Experiments were conducted to ascertain the biological effects of dosing

Table 2. Synopsis of Toxicology Procedures and Number of Animals*

Duration (htt) Animals Skin Intrinston (11) (11)	d	a si	Number of	Toxicolo	Toxicologic Evaluation ^c
(11) (11)	Group	Duration (hr)	Animals	Skin Irritation	Histopathology
HN or L) LL LL LL LL LL LL LL LL LL	Phase II ^d		(11)	(11)	(11)
1	Agent/Chloroform				
1	(10% HD, HN or L)				
14	5 µL	7	(2/11)	(2/11)	(2/11)
u.t. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1.	10 µL	7	(4/11)	(4/11)	(4/11)
1	50 µL	2	(2/11)	(2/11)	(2/11)
1	Neat HD (1 ,.L)	2	(4/11)	(4/11)	(4/11)
HCI3/t-BuOH 1 (5/11) (5/11) (5/11) HCI3/t-BuOH 1 (24) (24) (24) (20/24) 1 (4/24) (4/24) (4/24) (16/24)	Neat HD (1 µL)	-	(7/11)	(7/11)	(11/1)
HCl ₃ /t-BuOH) 1 (5/11) (5/11) (5/11) HCl ₃ /t-BuOH) 1 (24) (24) HN or L) HN or L) HA (4/24)	Oxidant/Solvent (20 μ L)				
124 (24) (24) (25) (26)	(DCDMH/CHCl ₃ /t-BuOH)	_	(5/11)	(5/11)	(5/11)
roform ^a HN or L) μL $\mu $	Phase III ⁰		(24)	(24)	(24)
1 (20/24) (20/24) 1 (4/24) (4/24) 1 (16/24) (16/24) 1 (8/24) (8/24) 1 (4/24) (4/24) 1 (4/24) (4/24) 1 (4/24) (4/24) 1 (4/24) (4/24) 1 (4/24) (4/24)	Agent/Chloroform				
1 (20/24) (20/24) 1 (4/24) (4/24) 1 (16/24) (16/24) 1 (8/24) (8/24) 1 (4/24) (4/24) 1 (4/24) (4/24) 1 (4/24) (4/24) 1 (4/24) (4/24)	(10% HD, HN or L)				
1 (4/24) (4/24) (16/24) (16/24) (16/24) (16/24) (16/24) (16/24) (16/24) (16/24) (16/24) (16/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24) (17/24)	5 µL	-	(20/24)	(20/24)	(20/24)
1 (16/24) (16/24) (16/24) (16/24) (8/24) (8/24) (4/24) (8/24) (8/24) (4/24) (4/24) (4/24) (8/24) (11 (8/24) (11/24) (11/24) (11/24) (11/24) (11/24) (11/24)	10 µL	-	(4/24)	(4/24)	(4/24)
1 (8/24) (8/24) (4/24) (4/24) (4/24) (4/24) (8/24) (8/24) (4/24) (4/24) (8/24) (8/24) (4/24) (4/24) (4/24)	Neat HD (1 µL)	-	(16/24)	(16/24)	(16/24)
1 (8/24) (8/24) (4/24) (4/24) (4/24) (4/24) (8/24) (8/24) (4/24) (4/24) (4/24) (4/24) (4/24) (4/24) (4/24)	Wastestreams				
1 (8/24) (8/24) (8/24) (4/24) (4/24) (4/24) (4/24) (4/24) (4/24) (4/24) (4/24) (8/24) (1 (8/24) (4/24) (4/24) (4/24)	"Archived" "Blue"			9	
1 (4/24) (4/24) (4/24) 1 (8/24) (4/24) (4/24) (4/24) (8/24) (8/24) (8/24) (4/24) (4/24)	25 µL	-	(8/24)	(8/24)	(8/24)
1 (8/24) (8/24) (4/24) (4/24) (4/24) (4/24) (8/24) (8/24) (4/24) (1 (8/24) (4/24) (4/24)	10 µL	_	(4/24)	(4/24)	(4/24)
1 (8/24) (8/24) 1 (4/24) (4/24) 1 (8/24) (8/24) 1 (4/24)	"Archived" "Red" h				
1 (4/24) (4/24) 1 (8/24) (8/24) 1 (4/24) (4/24)	25 µL	-	(8/24)	(8/24)	(8/24)
1 (8/24) (8/24) 1 (4/24) (4/24)	10 µL	-	(4/24)	(4/24)	(4/24)
1 (8/24) (8/24) 1 (4/24) (4/24)	"Archived" "Charcoal"				
1 (4/24)	25 µL	-	(8/24)	(8/24)	(8/24)
	10 uL	-	(4/24)	(4/24)	(4/24)

Synopsis of Toxicology Procedures and Number of Animals* (cont'd)

Treatment	Exposure	Number of	Toxicolo	Toxicologic Evaluation ^c
Group	Duration (hr)	Animais	Skin frritation	Histopathology
Wastestreams				
"Fresh" "Blue" (25 μ L) l	-	(8/24)	(8/24)	(8/24)
"Fresh" "Red" (25 μ L) †	-	(8/24)	(8/24)	(8/24)
"Fresh" "Charcoal" (25 μ L)	1	(4/24)	(4/24)	(4/24)

Toxicology studies comprised of two phases (Phase II and Phase III) of a multiphase effort: Phase II (dosing-ranging/optimization); Phase III (Vesicancy testing of wastestreams). The total number of animals on test was thirty five (Phase II studies (11); Phase III studies (24) æ

Multiple dosing sites per animal (refer to Fig 1). The number of animals per particular treatment ("test article"/dosage/exposure duration) and toxicologic evaluation is given as (#/#). 9

Toxicologic evaluation consisted of gross changes (erythema/edema) and light microscopic examination. Evaluation, (scoring of gross lesions) based on observations at 24-hr post-dosing. Following euthanasia, skin samples were taken and processed for microscopic examination. Ð

In Phase II studies, each animal was dosed dermally with neat HD, agent/CHCl, solution, and oxidant/solvent. Dosage of neat HD (1.0 μ L), dosage of agent/CHCl₃ solution (5, 10 and 50 μ L), and dosage of oxidant/solvent (20 μ L). "Test article" was allowed to remain in contact with the skin for either one or two hours. Dosages of "test article" rotated among skin exposure sites to control for differences in skin thickness.

Actual ampoules from CAIS kits were not used. Instead, "CAIS" (agent/chloroform solutions) were prepared from agent (HD, HN or L) stocks to the following specifications (10% HD, HN or L). <u>e</u>

In Phase III studies, each animal was dosed dermally with neat HD, agent/CHCI, solution, and wastestreams. Dosage of neat HD (1.0 μ L), dosages of agent/CHCl $_3$ solution (5 and 10 μ L), and wastestreams (10 and 25 μ L). Exposure duration was for one hour. Dosage of "test article" rotated among Chemical agents found in CAIS include HD, HN or L. Sulfur mustard (HD) was used as representative vesicant for the blistering agents. skin exposure sites to control for differences in skin thickness. E B

"Archived" "Blue" and "Red" wastestreams initially analyzed Oct 95 and re-analyzed and tested for vesicancy (Mar/Aug 961."Charcoal" wastestream initially analyzed Nov 95 and re-analyzed and tested for vesicancy in (Mar/Aug 96). Ξ

"Fresh" indicates that chemical analysis of wastestreams matched in time with bioassay.

volume and exposure duration, the uniformity and reproducibility of responses, and the skin-injurious effects of oxidant/solvent solution. Eleven animals were dermally dosed with neat HD (1 ul) and with 10 percent agent (HD, HN or L) in chloroform. Dosing volumes ranged from 5 to 50 ul, and exposure times were 1 or 2 hour durations. Five guinea-pigs were treated with oxidant/solvent solution. The exposed skin was examined 24-hr post-exposure for presence of gross and microscopic changes.

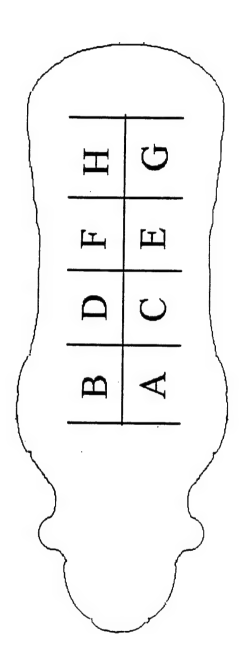
Phase III. This phase was designed to ascertain the vesicant potential of neutralized CAIS ("archived" and "fresh" wastestreams) and that of agent/chloroform solutions. The exposed skin was examined 24 hours after "test article" application for presence of skin-injurious effects (gross and microscopic).

2.4.2 Care and Treatment of Animals

A total of 35 male (approximately 200-350 g and 3 to 4 weeks of age upon receipt), euthymic hairless guinea-pigs (Cr1:IAF (HA)-hr BR), procured from Charles River Laboratories (Wilmington, MA; animals supplied from Portage, MI facility), were used in this study. Animals were quarantined and screened for general condition and health status, and were maintained in a program accredited by the Association for the Assessment and Accreditation of Laboratory Animal Care (AAALAC) International. Ear tags were applied to maintain positive identification, and animals were maintained between approximately 64 and 79 degrees F and 40 to 70 percent relative humidity with a 12-hr diurnal light cycle. Food and water were provided ad libitum and animals were housed individually in polycarbonate cages prior to exposure to "test article". Following treatment, animals were housed individually within a chemical fume hood during the 24-hr post-exposure period. Following recovery from anesthesia, animals were given food and water.

Animal Preparation and Dosing

Initially using 6 mg xylazine hydrochloride and 35 mg ketamine hydrochloride per kg of body weight given intramuscularly and increasing this to 13 mg xylazine and 87 mg ketamine/kg following the first day of dosing, anesthetized guinea-pigs were dosed topically on both sides of the dorsal midline with "test articles" (six to eight exposure sites/animal) - see Fig 1. Table 2 presents a synopsis of treatments, application volumes, and exposure durations. Approximately 24 hours after dosing, the animals were again anesthetized, sites evaluated for erythema and edema and lesion size, and animals then sacrificed with an inhalation anesthetic (halothane) overdose. Following euthanasia skin samples were collected and processed for histopathology.



2.5 Histopathologic Analysis

Following euthanasia, skin from the dosed sites was taken and placed in buffered formalin. After fixation, embedding, and sectioning, skin samples were stained with hematoxylin and eosin (H&E) and evaluated for histopathology. Histopathologic lesions (microblisters, epidermal necrosis, follicular necrosis, dermal necrosis, vascular necrosis, hemorrhage, and pustular epidermitis) were graded on a scale of 0-4, where 0 = normal, 1 = minimal, 2 = intermediate, 3 = moderate, 4 = severe. Definitions for scoring histopathology and the criteria for grading severity of lesions are summarized in Table 3. The grading of microblister formation is highlighted in Table 4.

2.6 DATA ANALYSIS

For chemistry data generated in Phase I, means and standard deviations of responses of each control standard were determined to calculate both the inter- and intra- variability of the analytical method. Calibration performance characteristics for each analyte, such as slope and standard error of the slope, R² (measure of fit about the regression line), method detection limits, and quantitation limits were calculated.

For Phase III data (vesicating assessment of wastestreams), statistical hypothesis tests were conducted at the 5 percent significance level to determine whether or not the neutralization process reduced the vesicating property of agents contained in CAIS. For each CAIS sample, the incidence of microblisters at sites treated with CAIS agent(s) were compared to those of contralateral sites treated with the wastestream. Although incidence of microblisters was the primary endpoint for evaluating the efficacy of each neutralization process, analyses were also conducted on other indices of skin injury (gross and microscopic). To accommodate the intra-animal correlation of multiple measurements made on the same animal, McNemar's test was used to analyze quantal data (Agresti, 1990). Analysis of variance (ANOVA) models, that include random effects for animal, were fitted to continuous data. If data were not approximately normal, ANOVA were conducted on transformed data, or nonparametric or categorical methods of analysis were performed.

TABLE 3. DEFINITIONS USED IN HISTOPATHOLOGIC EVALUATIONS AND AN EXPLANATION OF THE GRADING OF LESION SEVERITY

Microbilister	Loss of epidermal basal cell attachment to the underlying basement membrane of at least two adjacent cells. The loss of attachment creates a space which may appear empty, full of proteinaceous fluid, or filled with neutrophils. One or a few isolated small areas of detachment is graded 1, minimal. Many such areas of detachment, or several larger (10 or more contiguous cells) areas of detachment is graded 2, intermediate. When half or more of the epidermis in the tissue section is detached from the dermis, it is graded 3, moderate. Such lesions typically have a much larger space between the basal cells and the dermis. When nearly all of the epidermis is separated from the dermis, it is graded 4, severe. In such situations, there are usually focal, point attachments, so the entire epidermis is not lifted along the full width of the section.
Epidermal necrosis	The epidermal cells exhibit cytoplasmic eosinophilia, nuclear loss or pyknosis, and are generally shrunken. If only individual cells are affected, it is graded 1 (these are generally isolated basal cells). If small areas are affected, with normal areas in close proximity, it is graded 2. If the epidermis exhibits cell death in a full-thickness (all layers of epidermis) pattern, and affects half or more of the skin section, it is graded 3. If the epidermis is virtually entirely necrotic, it is graded 4. Severe ulcers assume that the epidermis is necrotic.
Follicular necrosis	If isolated epithelial cells of the hair follicles exhibit eosinophilia or pyknosis, it is graded 1. If clusters of adjacent cells within follicles are dead, it is graded 2. If cells of half or more of a particular hair follicle are dead, it is graded 3. Grade 4 lesions have complete necrosis of the follicular cpithelium underlying much of the epidermal lesion area. This indicates that the agent has penetrated deeply.
Dermal necrosis	Loss of collagen fiber integrity, evidenced by pale eosinophilic staining and homogeneous appearance, indicates necrosis of dermal fibers. With only isolated areas, it is graded 1. Multiple areas are graded 2. Necrosis of most of the superficial dermal collagen in the lesion area is graded 3. A grade 4 lesion requires deep (to the base of the associated adnexa) dermal necrosis.
Vascular necrosis	Loss of integrity of a medium to large blood vessel is vascular necrosis. Grading depends upon the number of vessels affected and the severity. Partial necrosis of one vessel is graded 1 to 2. Complete necrosis of a vessel is graded 3; multiple such lesions are graded 4.
Hemorrhage	Extravasated erythrocytes is hemorrhage. A few isolated foci is graded 1. Multiple, common foci is graded 2. Large pools of blood is graded 3. A grade 4 lesion requires a massive area of blood pooling and the displacement of large areas of dermal collagen.
Pustular epidermitis	Collections of neutrophils in the epidermis proper is graded by extent; one or two small foci is graded 1; three or more small foci is graded 2; one or more large foci is graded 3; a grade 4 lesion would indicate massive infiltration of the entire epidermis by neutrophils

TABLE 4. DEFINITION OF DEGREES OF SEVERITY USED FOR HISTOPATHOLOGIC EVALUATION OF VESICATION (MICROBLISTER FORMATION*)

Lesion Characteristic	Degree of Severity
No lesion (unaffected) One or a few isolated areas of detachment Many small areas of detachment or several larger areas of detachment >50% of the epidermis in tissue section is detached from the dermis (much larger space between basal cells and dermis) Nearly all the epidermis is separated from the dermis	0 (normal) 1 (minimal) 2 (intermediate) 3 (moderate) 4 (severe)

^a Microblister: loss of epidermal basal cell attachment to underlying basement membrane of at least two adjacent cells. Loss of attachment creates a space.

3. Results

3.1 Chemistry

Nitrogen mustard, sulfur mustard, and lewisite are components of CAIS that were chemically neutralized ("detoxified") on reaction with treatment reagent (1, 3-dichloro-5, 5dimethylhydantoin). The selection of a particular process chemistry (designated as "Blue", "Red", or "Charcoal" process) was dependent on whether the agent was neat material (HD), in solution (agent in chloroform), or adsorbed on charcoal, The DCDMH-mediated neutralization of sulfur mustard resulted in HD concentrations below 50 ppm in "Blue" process wastestream (product solution). Chemical treatment resulted in the conversion of sulfur mustard to HD sulfoxide degradation products (Further oxidation to sulfone was also a possibility under conditions via neutralization by DCDMH). Secondary reactions (i.e. elimination/substitution) also occurred that produced chlorinated and vinyl sulfoxides. The neutralization reaction between oxidant and CAIS containing agent (HD, HN or L in chloroform - "Red" process) resulted in complex product solutions containing various products/by-products and residual amounts of unreacted agent. The process chemistry for neutralization of CAIS components containing agent (HD, HN or L) on charcoal ("Charcoal" process), also resulted in the formation of complex product solutions. Residual amounts of agent were detected. Details pertaining to the process chemistry and analyses have been reported in detail (Olson et al, 1997 and Lucas, 1997).

The "archived" wastestreams were additionally analyzed for product/by-product composition. HD sulfoxide and other degradation products resultant from secondary reactions (e.g. elimination, substitution) were detected in wastestream samples. HD sulfone and/or its vinyl containing derivatives, which are known vesicants, were not detected in the "Blue" process wastestream. Product analyses did not reveal HD sulfone or vinyl/divinyl analogs in the product solution obtained from the chemical neutralization of CAIS containing agent in chloroform ("Red" process). Product characterization of the "Charcoal" wastestream did not reveal HD sulfone; however, multichlorinated vinyl containing derivatives (non-vesicant) were present in the product solution.

3.2 Dermal Effects

3.2.1 Gross Pathologic Findings

Phase II. All skin exposures to HD and agent/chloroform solutions containing 10 percent HD, HN or L resulted in gross skin lesions consisting of well-defined areas of edema and erythema of moderate to severe intensity. In some instances, large areas of ulceration with complete loss of the covering epidermis was evident. The skin-injurant effects of HN and L were comparable to that produced by HD (refer to Table 5 and Appendix A). The skin-injurious effect of oxidant/solvent solution was minimal gross lesions (refer to Table 5 and Appendix A).

Phase III. The cutaneous injury (non-vesicant) effects after one hour exposure to HD, agent/CHC1₃, or CAIS wastestreams ("archived" and "fresh") were evaluated and are summarized in Table 6. Individual gross pathology data are presented in Appendix A. All agent-dosed sites demonstrated gross lesions. Wastestream-induced dermal injury resulted in mild to moderate degrees of erythema and edema.

3.2.2 Histopathologic Findings

Phase II. Two hour dermal exposures of animals to neat HD (1 μ L) and to various doses (5 - 50 μ L) of agent/chloroform solutions containing 10 percent HD, HN or L resulted in microblister formation of intermediate to severe intensity - refer to Table 7. Incidence of histopathologic changes are summarized in Table 8. In some animals, large areas of ulceration with loss of epidermis prevented the occurrence of microblisters. Individual animal histopathology data are presented in Appendix B. Based on the outcome of the two-hour exposure studies, other guinea pigs were dosed with 5 and 10 μ L volumes of 10 percent agent in chloroform solutions and with neat HD (1 μ L) at an exposure duration of one hour. Microblister formation was evident at all sites, unless occurrence was precluded by development of an ulcer, and ranged in severity from moderate to severe. The application of 5 μ L of 10 percent agent/chloroform solution resulted in microblisters of at least intermediate severity. Refer to Table 7 for incidence/response summary and Appendix B for individual histopathologic findings. The oxidant/solvent system was also evaluated for skin effects. Animals treated with oxidant/solvent solution did not manifest dermal lesions other than minimal inflammatory cell infiltration - refer to Table 8 and Appendix B.

Phase III. Twenty-four animals comprising Phase III of the study were treated with "neutralized" CAIS to ascertain the vesicating potential of chemically degraded CAIS. Incidence/response data related to microvesication are summarized in Tables 9, 10, and 11. A

TABLE 5. PHASE II - SKIN REACTION (ERYTHEMA AND EDEMA) FOLLOWING EXPOSURE TO HD, AGENT/CHCL, SOLUTIONS, AND OXIDANT/SOLVENT SOLUTION

Experiment Date/ Animal ID	Test Article	Dose Volume (µL)	Time to Decontamination (hr)	No.01 Animals Tested	Erythema Score, Mean	Edema Score, Mean
	10% L/CHCl ₃	10	2	2	3.0	3.0
	10% L/CHCl ₃	50	2	2	3.0	3.0
	10% HN/CHCl ₃	10	2	2	2.0	2.0
. 02/19/96	10% HN/CHCl ₃	50	2	2	2.0	2.0
(301, 305)	10% HD/CHCl ₃	10	2	2	2.0	2.0
	10% HD/CHCl ₃	50	2	2	2.5	2.0
	Neat HD	1	2	2	2.0	2.0
	10% L/CHCl ₃	5	2	2	2.5	3.0
	10% L/CHCl ₃	10	2	2	2.5	3.0
	10% HN/CHCl ₃	5	2	2	2.0	2.0
02/21/96 (306, 309)	10% HN/CHCl ₃	10	2	2	2.0	2.5
	10% HD/CHCl ₃	5	2	2	3.0	3.0
	10% HD/CHCl ₃	10	2	2	2.0	2.0
	Neat HD	1	2	2	2.0	2.5
	10% L/CHCl ₃	5	1	2	3.0	3.0
	10% L/CHCl ₃	10	1	2	3.0	3.0
	10% HN/CHCl ₃	5	1	2	2.0	2.5
02/27/96 (312, 316)	10% HN/CHCl ₃	10	1	2	2.0	2.0
(312, 310)	10% HD/CHCl ₃	5	1	2	3.0	2.0
	10% HD/CHCl ₃	10	1	2	2.5	2.0
	Neat HD	1	1	2	3.0	2.5
	10% L/CHCl ₃	5	1	5	3.0	2.8
03/05/96	10% HN/CHCl ₃	5	1	5	1.8	2.0
(311, 313, 315, 317,	10% HD/CHCl ₃	5	1	5	2.4	2.4
324)	Neutralizing Solution	20	1	5	0.0	1.0
	Neat HD	1	1	5	2.4	2.6

TABLE 6. PHASE III. SKIN REACTION (ERYTHEMA AND EDEMA) FOLLOWING EXPOSURE TO HD, AGENT/CHCI, SOLUTION OR CAIS WASTESTREAMS

Date,		Dose	No of						
Source of		Volume	Animals	Erythen	Erythema Score	Edemi	Edema Score	Lesion Area (mm²)	.a (mm²)
Wastestream	Test Article	(III)	Tested	Mean	S.D.	Mean	S.D.	Mean	us
	10% L/CHCl,	5	∞.	2.9	0.3	3.0	0.0	P \$6	22.2
03/13/96.	10% HN/CHCI,	2	&	2.0	6.0	2.0	0.5	109	13.6
03/21/96	10% HD/CHCl,	~	∞	2.6	0.7	2.1	0.8	107.9	35.8
1 A	"Red" Wastestream	25	&	1.1 26,0	0.3	1.8 *	0.5	237 4 de.f	71.3
Wastestreams	"Blue" Wastestream	25	80	1.9 %	8.0	1.6 %	0.5	236 5de.f	77.6
	"Charcoal" Wastestream	25	8	0.4 ab.c	0.2	0.4 abe	0.2	132 %	050
	Neat HD	1	œ	2.8	0.5	2.9	0.3	180.2	53.1
	10% L/CHCI,	5	8	3.0	0.0	3.0	0.0	1560	1.69
06/20/96,	10% HN/CHCI,	\$	8	1.9	0.3	2.1	0.3	82.0	30.0
06/56/96	10% HD/CHCI,	5	∞	2.4	0.5	2.3	0.5	94.9	18.4
"Fresh"	"Red" Wastestream	25	8	0.3 4.6.0	0.3	0.3 abe	0.3	46.2	\$2.3
Wastestreams	"Blue" Wastestream	25	00	1.8 20	0.5	1.6 abc	0.5	220 6 de,f	42.0
	Neat HD	1	8	2.5	0.5	2.4	0.5	126.8	32.6
	10% L/CHCl,	10	4	3.0	0.0	2.8	0.5	212.6	35.6
08/13/96	10% HN/CHCI,	10	4	2.5	9.0	2.3	0.5	155.1	21.4
	10% HD/CHCI,	10	4	2.3	1.0	2.3	0.5	178.7	34.9
Wastestreams	"Red" Wastestream	10	4	1.1 abc	9.0	0.8 a.b.c	1.0	121.140	41.8
	"Blue" Wastestream	10	4	1.3 ab.c	0.5	1.0 4.6.0	0.0	142.4*	42.3
	"Charcoal" Wastestream	10	4	0.4 a.b.c	0.2	0.0 26.0	0.0	69.34b.c	23.0
08/23/96	10% L/CHCI,	5	4	3.0	0.0	2.8	0.5	113.9	33.1
:	10% HN/CHCI,	5	4	1.5	9.0	2.0	0.8	91.3	19.6
"Fresh" Wastestream	10% HD/CHCl,	S	4	3.0	0.0	3.0	0.00	89.5	26.2
	"Charcoal" Wastestream	25	4	0.0 abc	0.0	0.0 4.6.0	0.0	0.046.0	0.0

Note: All times to decontamination were I hr.

a Mean is significantly less than that observed on sites treated with L. b Mean is significantly less than that observed on sites treated with HN.

Mean is significantly less than that observed on sites treated with HD.

Mean is significantly greater than that observed on sites treated with L. Mean is significantly greater than that observed on sites treated with HN.

Mean is significantly greater than that observed on sites treated with HD.

TABLE 7. PHASE II. VESICATION (MICROBLISTER FORMATION) IN HAIRLESS GUINEA PIGS FOLLOWING DERMAL EXPOSURE TO HD, AGENT/CHCI, SOLUTIONS, OR NEUTRALIZING SOLUTION (DCDMH/CHCI,/t-BuOH)^a

				Micro	Microblister Severity (0-4)	(0-4)				
Treatment b	Animal									Mean
Group (2 hour)	No.	301	305	306	309				Response	Severity
Neat HD (1 µL)		2	2	0	г				4/4	2.0
50 uL		7	7						2/2	2.0
10 µL		2	٣	က	m				4/4	2.8
SμL				۰,	0				1/2	0.5
10% HN/CHCI,										
50 µL		7	7	,					2/2	2.0
10 µL		7	7	2 c	4				4/4	2.5
				4	4				2/2	4.0
10% L/CHCl ₃										
50 µL		4	m						2/2	3.5
10 µL		3	æ	3	4				4/4	3.3
SµL				4	4				2/2	4.0
Treatment	Animal									Mean
Group (1 hour)	No.	312	316	311	313	315	317	324	Response	Severity
Neat HD (1 µL)		3	3	3	2	2	7	33	LIL	2.6
10% HD/CHCl,										,
10 µL		m	m						2/2	3.0
SµL		3	e	2	E	e	2	4	L/L	2.9
10% HN/CHCI,										
10 µL		4	, ,						2/2	3.5
SµL		3	4	æ	4	2	3	4	LIL	3.3
10% L/CHCI,									!	•
10 nL		m	4						2/2	5.5
5μL		m	4	e	4	4	7	4	<i>L/L</i>	3.4
DCDMH/CHCl ₃ /				0	0	0	0	0	9/2	0
(20 uL)										

a At 24 hr after dosing, animals were evaluated for skin injury, sacrificed, and skin samples taken and prepared for histopathology.
 b Exposure duration 2 hr.
 c Ulceration at dosing site may have obscured evidence of microvesication
 d Exposure duration 1hr.

TABLE 8. PHASE II. SUMMARY OF HISTOPATHOLOGY RESULTS

Experiment		Dose	Time to					Numb	Number of Animals with Sign	ith Sign		
Date/ Animal ID	Test Article	Volume (µL)	Decon. (hr)	No.of Animals	No. of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular
	10% L/CHCl3	10	2	2	2	2	2	2	0	0		
	10% L/CHCl,	50	2	2	2	2	2	2	0	0	-	0
02/19/96	10% HN/CHCl3	10	2	2	2	2	2	2	0	0	0	0
100	10% HN/CHCI3	50	2	2	2	2	2	2	0	0	0	0
(301, 305)	10% HD/CHCI,	10	2	2	2	2	2	2	0	0	_	0
	10% HD/CHCI,	50	2	2	2	2	2	2	0	0	_	0
	Neat HD	1	2	2	2	2	2	2	0	0	0	0
	10% L/CHCI3	5	2	2	2	2	2	2	0	-	_	0
	10% L/CHCl3	10	2	2	2	2	2	2	0	-	-	0
02/21/96	10% HN/CHCI,	5	2	2	2	2	2	2	1	0	0	0
	10% HN/CHCI,	10	2	2	2	2	2	2	-	-	0	0
(306, 309)	10% HD/CHCI,	5	2	2	2	1	2	2	0	2	0	0
	10% HD/CHCl3	10	2	2	2	2	2	2	1	-	0	0
	Neat HD	-	2	2 .	2	2	2	2	0	-	0	0

TABLE 8. PHASE II. SUMMARY OF HISTOPATHOLOGY RESULTS (CONT'D.)

Exneriment		Dose	Time to					Num	Number of Animals with Sign	ith Sign		
Date/ Animal ID	Test Article	Volume (µL)	Decon. (Hr)	No. Of Animals	No. of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular Necrosis
	10% L/CHCI,	5	-	2	2	2	2	2	0	0	2	0
	10% L/CHCl,	10	1	2	2	2	2	2	0	0	2	0
20110	10% HN/CHCl3	\$	1	2	2	2	2	2	2	0	0	0
06/17/70	10% HN/CHCl ₃	01	-	2	2	2	2	2	2	0	0	0
(312, 316)	10% HD/CHCl ₃	5	-	2	2	2	2	2	1	-	0	0
	10% HD/CHCl ₃	10	-	2	2	2	2	2		0	-	0
	Neat HD	-	-	2	2	2	2	2	1	0	0	0
	10% L/CHCl,	5	1	8	5	\$	5	5	0	-	5	0
03/02/96	10% HN/CHCi	5	-	5	\$	5	5	5	4	-	-	0
(311 313	10% HD/CHCl ₃	5	1	5	5	5	5	2	-	3	2	0
315, 317, 315, 317, 324)	Neutralizing Solution	20	-	\$	20	0	0	. 0	0	0	0	0
	Neat HD	1	-	5	5	5	\$	2	0	0		0

summary of histopathologic changes, including vesication, is presented in Tables 12 and 13. Individual histopathology data appear in Appendix B. Eight animals were dosed with "archived" wastestreams, agent/chloroform solutions, and neat HD. Guinea-pigs dosed with HD and agent/chloroform solutions demonstrated at least minimal microvesication along with consistent, marked epidermal and follicular necrosis. The "Blue" process wastestream ("archived"; 25 μ L application) resulted in intermediate to severe microblisters and severe epidermal necrosis at all sites dosed (refer to Tables 9 and 13 and Appendix B). The impression of the pathologist reading the slides was that lesions did not appear to be "basal cell specific", as chemical blistering agents appear to cause, nor did the lesions resulting from application of the "Blue" wastestream penetrate deeply enough to cause severe necrosis in the follicular epithelium. A photomicrograph representative of the morphologic changes observed following treatment with a vesicant is shown in Figure 2a, and one demonstrating the appearance of normal hairless guinea pig epidermis is shown in Figure 2b. The morphologic changes seen consist of ballooning degeneration and loss of epidermal basal cell attachment to the underlying basement membrane. Neither "Red" nor "Charcoal" process wastestreams ("archived"; 25 µL application) produced microblisters (Tables 9 and 12). The "Red" process wastestream produced only minimal pustular epidermitis or minimal epidermal necrosis (refer to Table 12 and Appendix B). The "Charcoal" process wastestream ("archived"; 25 μ L application) killed some surface epithelial cells (minimal to intermediate epidermal necrosis) but did not penetrate to basal cells - refer to Table 12 and Appendix B. Four guinea pigs were dosed with 10 µL of "Blue", "Red", and "Charcoal" process wastestreams ("archived") and evaluated for dermal effect. The "Blue" process wastestream induced microblisters whereas the "Red" and "Charcoal" process wastestreams did not elicit microblister formation. The findings are highlighted in Table 10. Histopathology findings are summarized in Tables 12 and 13, and individual histopathology data are presented in Appendix B.

"Fresh" wastestream-induced skin effects were also evaluated. Data on microvesication are presented in Tables 11, 12 and 13, and other histopathologic skin effects data are given in Tables 12 and 13. Individual animal histopathology results are presented in Appendix B. All agent-dosed sites (neat HD and agent/chloroform solutions) and all "Blue" process wastestream sites demonstrated histopathologic lesions including microvesication. In "fresh" "Red" process wastestream-dosed animals, minimal to no lesions were seen on histopathologic examination. One "Red" process wastestream site in one animal demonstrated histopathology, including minimal microvesication; however, this lesion was incompatible with what had been noted previously. The "Charcoal" process wastestream did not produce microblisters and none of the sites demonstrated histopathology graded more than minimal.

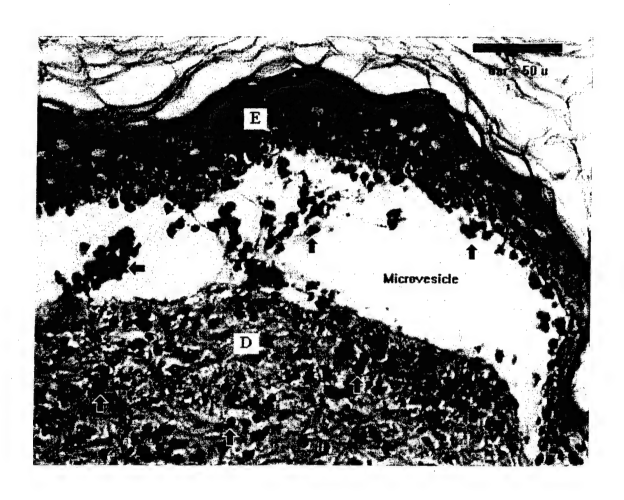


Fig. 2a. Typical microblister in a hairless guinea pig 24 hours after exposure to vesicant. Epidermis (E) is eosinophilic and shrunken due to necrotic epithelium; dermis (D) is also necrotic and contains an infiltrate of polymorphonuclear cells (arrows), as does the microblister cavity (microvesicle).

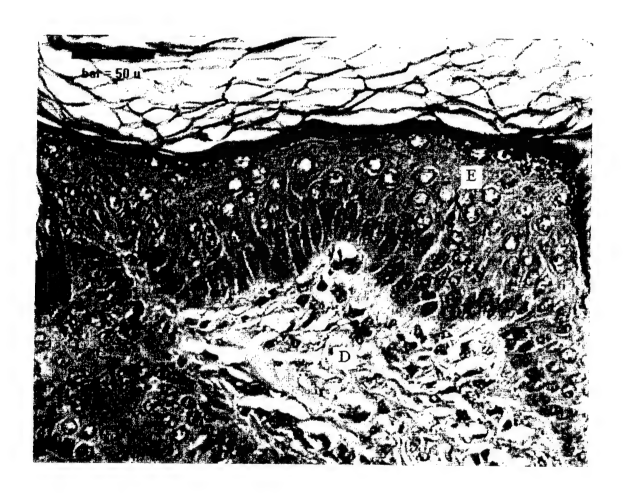


Fig. 2b. Normal skin from a hairless guinea pig. Epidermis (E) and dermis (D) are visible. Note differences in appearance from the necrotic tissue depicted in Fig. 2a. Magnification of both skin photomicrographs is the same.

PHASE III. VESICATION (MICROBLISTER FORMATION) IN HAIRLESS GUINEA PIGS FOLLOWING EXPOSURE TO "ARCHIVED" RRS WASTESTREAMS, AGENT/CHCI, SOLUTIONS, OR NEAT SULFUR MUSTARD (HD) 4,b TABLE 9.

- ⁻2

						Microblister Severity (0-4)	ter Severi	ty (0-4)			
											Mean
Treatment Group	Animal No.	la! 494	496	497	499	310	491	493	498	Response	Severity Score
Neat HD (1[.)		3	2	2	3	1	2	2		8/8	2.0
10% HD/CHCl,	(5 µL)	_	0	2	4	2		-	3	2/8	1.8
10% HN/CHCI,	(5 µL)	3	0	-	4	4	4	4	2	8/L	2.8
10% L/CHCI,	(5 µL)	4	-	4	3	_	4	7	3	8/8	2.8
"Blue" wastestream	$(25 \mu L)$	2	4	2	2	7	3	4	3	8/8	2.8
"Red" wastestream	$(25 \mu L)$	0	0	0	0	0	0	0	0	8/0	0
"Charcoal" wastestream (25 μ L)	n (25 µL)	0	0	0	0	0	0	0	0	8/0	0

a Each animal was dosed percutaneously (1 hr exposure) with neat HD, agent/ CHCl3 solution, and "archived" wastestreams. Sites were evaluated visually at about 24 hr after dosing, and the animals then sacrificed and skin samples taken and prepared for histopathologic evaluation.

b Dosing volumes of HD and agent/CHCl₁ solutions, as well as the duration of exposure, were based on preliminary tests. Dosing volumes of wastestreams were based upon approximate ratio of neutralization solution volume to volume of agent treated.

c Wastestreams were generated from the reaction of DCDMH (oxidant) with heat HD ("Blue" process), with 10% HD, HN or L in CHCl, ("Red" process), or with HD, HN, or L on charcoal ("Charcoal" process).

TABLE 10. PHASE III. MICROBLISTER FORMATION IN HAIRLESS GUINEA PIGS FOLLOWING EXPOSURE TO EQUAL VOLUMES OF "ARCHIVED" RRS WASTESTREAMS OR AGENT/CHCI, SOLUTIONS **

					Microbli	Microblister Severity (0-4)	ity (0-4)	
Treatment Group		Animal No.	383	385	389	400	Response	Mean Severity Score
10% HD/CHCl,	$(10 \mu \text{L})$		2	ю	7	æ	4/4	2.5
10% HN/CHCI,	$(10\mu\text{L})$		ю	4	m	4	4/4	3.5
10% L/CHCI,	$(10 \mu L)$		ю	4	2	, E	4/4	3.0
"Blue" wastestream	$(10 \mu L)$		0	-	2	m	3/4	1.5
"Red" wastestream	$(10 \mu \text{L})$		0	0	0	0	0/4	0
"Charcoal" wastestream	$(10 \mu \text{L})$		0	0	0	0	0/4	0

a Each animal was dosed dermally (1 hr exposure) with agent/CHCl₃ solutions and wastestreams. Sites were evaluated visually at about 24 hr after dosing, and the animals then sacrificed and skin samples taken and prepared for histolopathologic evaluation.

b Wastestreams (product solutions) generated from reaction of oxidant (DCDMH) with HD - "Blue"; 10% HD, HN or L in CHCl3 - "Red"; HD, HN or L on charcoal - "Charcoal".

TABLE 11. PHASE III. MICROBLISTER FORMATION IN HAIRLESS GUINEA PIGS FOLLOWING EXPOSURE TO "FRESH" RRS WASTESTREAMS, AGENT/CHCI, SOLUTIONS, OR NEAT HD ^{a,b}

Microblister Severity (0-4)

Treatment Group		Animal No.	339	341	342	346	340	345	351	352	Response	Mean Severity Score
Neat HD	(i μL)		2	7	m	. 7	0	-	-	7	8/L	1.6
10% HD/CHCl,	$(5 \mu L)$		3	2	٣	7	2	2	-	_	8/8	2.0
10% HN/CHCI,	$(5 \mu L)$		ю	7	4	4	60	-	-	7	8/8	2.5
10% L/CHCI,	$(5 \mu L)$		4	m	ю	2	ю	3	4	3	8/8	3.1
"Blue" wastestream	$(25 \mu\text{L})$		2.5	_	7	-	60	1.5	7	1.5	8/8	1.8
"Red" wastestream	$(25 \mu \text{L})$		0	0	0.5 ^d	0	0	0	0	0	1/8	0
Treatment Group		Animal No.	379	380	387	388					Response	Mean Severity Score
10% HD/CHCl ₃	(5 µL)		7	E	ω	4					4/4	3.0
10% HN/CHCl ₃	$(5 \mu L)$		Э	4	2	3					4/4	3.0
10% L/CHCl,	$(5 \mu L)$		4	4	4	4					4/4	4.0
"Charcoal" wastestream	$(25 \mu L)$		0	0	0	0					0/4	0

Each animal was exposed dermally for 1 hr to "test article" (neat HD and/or agent/CHCl, solution, and wastestreams). At 24 hr after dosing, animals were evaluated for gross skin injury and then sacrificed and skin samples taken and prepared for histopathologic evaluation.

Dosing volumes and duration of exposure were determined from preliminary testing. Dosing volume of wastestreams was selected on the basis of approximate neutralization solution volume to volume to volume of agent. Wastestreams were generated via the reaction of DCDMH with neat HD - "Blue"; HD, HN, or L in CHCl₃ -"Red"; and HD, HN, or L on charcoal - "Charcoal".

: Mean value for the two sites dosed with each wastestream on each animal.

d Could be due to adjacent HD-treated site.

e Three sites on each animal were dosed with "Charcoal" wastestream and no sites exhibited microblisters.

TABLE 12. PHASE III. SUMMARY OF HISTOPATHOLOGY RESULTS

2.2

Date,		Dose					Number	Number of Animals with Sign	h Sign		
Source of Wastestream	Agent / Compound	Volume (µL)	No. Of Animals	No.of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular
	J	5	8	8	8	∞	8	0	9	2	0
03/13/06	H.	5	80	80	7.	∞	80	3	4	0	0
03/21/96	HD	5	80	8	7.	•	∞	3	7	0	0
	"Red" Wastestream	25	80	8	0 b.c.d	p'o'q I	p'c'q ()	2	p.d 0	0	0
"Archived" Wastestreams	"Blue" Wastestream	25	80	8	8	∞	p'c'q l	-	1 q	0	0
	"Charcoal" Wastestream	25	80	8	0 b.c.d	9	p'c'q 0	2	p.4 0	0	0
	Neat HD	1	8	8	8	8	8	0	7	_	0
	Г	5	•	80	8	8	8	1	3	2	0
06/20/96,	뫂	5		8	8	∞	8	2	5	2	0
06/26/96	HD	5	•	∞	8	∞	8	2	5	8	0
"Fresh"	"Red" Wastestream	25	•	16	l b,c,d	2 b,c,d	p'o'q l	1	0	0	0
Wastestreams	"Blue" Wastestream	25	•	91	8	80	<i>L</i>	2	-	0	0
	Neat HD	-	•	8	7	8	8	0	5	S	0
	L	10	4	4	4	4	4	0	0	4	_
08/13/96	NH.	10	4	. 4	4	4	4	4	-	2	0
	HD	10	4	. 4	4	4	4	1	0	2	0
"Archived"	"Red" Wastestream	10	4	4	0	0	0	0	0	0	0
w dolcou caliis	"Blue" Wastestream	10	4	4	3	3	7	1	0	-	0
	Charcoal Wastestream	10	4	4	0	1	0	1	0	0	0
96/66/80	L	2	4	4	4	4	4	0	-	4	0
	HN	5	4	4	4	4	4	1	-	-	0
"Fresh"	HD	S	4	4	4	4	4	0	1	2	0
w astesticanis	"Charcoal" Wastestream	25	4	12	0	4	4	1	0	0	0

vote: All times to decontamination were 1 hr.

a Marked ulceration at the dosing site on animal number 496 obscured any evidence of microvesication.

b Incidence of sign was significantly less than that for sites dosed with L using McNemar's Test and a significance level of p=0.05.

c Incidence of sign was significantly less than that for sites dosed with HN using McNemar's Test and a significance level of p=0.05. d Incidence of sign was significantly less than that for sites dosed with HD using McNemar's Test and a significance level of p=0.05.

e Agent (L, HN, HD) at a concentration of 10% in chloroform.

TABLE 13. PHASE III. SUMMARY OF INTERMEDIATE TO SEVERE HISTOPATHOLOGY RESULTS

Description Agent of Notice of Notic	t of		200	W 1			Number	f Animals wi	Number of Animals with Sign Rated Intermediate to Severe	ntermediate	to Severe	
HN S S S S S S S S S	Source of Wastestream		Volume (µL)	No. Of Animals	No.of Sites	Micro- blister	Epidermal Necrosis	Follicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhage	Vascular Necrosis
HIV 5 8 8 8 6 6 9 9 9 9 9 9 9 9 9 9 9 9 9 9 9		7	5	8	80	-9	8	8	0	9	2	0
HD - 5 8 8 4*		呈	2	8	80	•9	8	œ	0	3	0.	0
"Red" Wastestream 25 8 8 0 hod 0 hod <t< td=""><td>03/13/96,</td><td></td><td>5</td><td>8</td><td>80</td><td>4.</td><td>æ</td><td>8</td><td>0</td><td>7</td><td>0</td><td>0</td></t<>	03/13/96,		5	8	80	4.	æ	8	0	7	0	0
"Glue" Wastestream 25 8 8 6 9 % 4 0 0 % 4 0 0 % 4 0 <t< td=""><td>03/21/30</td><td>"Red" Wastestream</td><td>25</td><td>00</td><td>80</td><td>o'q 0</td><td>p'c'q 0</td><td>p°c°q 0</td><td>0</td><td>p.4 0</td><td>0</td><td>0</td></t<>	03/21/30	"Red" Wastestream	25	00	80	o'q 0	p'c'q 0	p°c°q 0	0	p.4 0	0	0
"Charcoal" Wastestream 25 8 8 0 bts d	"Archived"	"Blue" Wastestream	25	•	&	æ	80	p'o'q0	0	p'q 0	0	0
Neat ID 1 8 6* 8 6* 8 6* 0 6 0 6 0 6 0 6 0 6 0 6 0 6 0 6 0 6 0 6 0 6 0 6 0 3 1 6 0 0 3 1 2 6 0 3 1 2 8 6* 8 6* 8 0 3 1 0 0 3 1 0 <t< td=""><td>Wastestreams</td><td>"Charcoal" Wastestream</td><td>25</td><td>••</td><td>8</td><td>o'q 0</td><td>2 b.c.d</td><td>0 p.c.d</td><td>0</td><td>P:q 0</td><td>0</td><td>0</td></t<>	Wastestreams	"Charcoal" Wastestream	25	••	8	o'q 0	2 b.c.d	0 p.c.d	0	P:q 0	0	0
HN 5 8 8 6 6 8 8 8 6 0 3 5 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6 6		Neat HD	-	•	8	•9	8	80	0	9	0	0
HN 5 8 6* 8 6* 8 6* 9 3 1 0 HD 5 8 6* 8 6* 8 6* 9 7 9 7 7 9 7 9 7 7 7 8 2 bcd 0 <		J	5	8	∞	&	8	œ	0	3	2	0
HD 5 8 6** 8 6** 8 6** 8 6** 8 6** 8 6** 6** 8 6** 9** 9 0 5 9 0	06/20/96.	NH.	5	∞	•	.9	8	∞	0	3	1	0
"Red" Wastestream 25 8 16 0 beed 1 bed 1 bed 0 <	06/26/96	HD	8	∞	00	.9	8	∞	0	2	2	0
"Blue" Wastestream 25 8 16 7 8 2 bcd 0 1 0 Neat HD 1 8 8 5 8 8 0 1 0 L 10 4 4 4 4 4 4 0 0 0 4 6 0 <t< td=""><td>"Erech"</td><td>"Red" Wastestream</td><td>25</td><td>oo</td><td>91</td><td>p'c'q 0</td><td>1 b,c,d</td><td>l b.c.d</td><td>0</td><td>0</td><td>0</td><td>0</td></t<>	"Erech"	"Red" Wastestream	25	o o	91	p'c'q 0	1 b,c,d	l b.c.d	0	0	0	0
Neat HD 1 8 8 5 8 8 1 6 4 </td <td>Wastestreams</td> <td>"Blue" Wastestream</td> <td>25</td> <td>∞</td> <td>91</td> <td>7</td> <td>8</td> <td>2 b,c,d</td> <td>0</td> <td>-</td> <td>0</td> <td>0</td>	Wastestreams	"Blue" Wastestream	25	∞	91	7	8	2 b,c,d	0	-	0	0
L 10 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 4 6 0		Neat HD	-	•	∞	5	8	∞	0	2	1	0
HIN 10 4 4 4 4 4 4 6 0 0 0 0 0 0 1 1 4 4 4 4 4 4 4 4 4 4 0		Γ	10	4	4	4	4	4	0	0	4	0
HD 10 4 4 4 4 4 4 6 10 10 4 4 4 6 10 0 <td></td> <td>HN</td> <td>10</td> <td>4</td> <td>4</td> <td>4</td> <td>4</td> <td>4</td> <td>0</td> <td>0</td> <td>0</td> <td>0</td>		HN	10	4	4	4	4	4	0	0	0	0
"Red" Wastestream 10 4 4 0 0 0 0 0 0 0 "Blue" Wastestream 10 4 4 2 2 0	08/13/96	HD	01	4	4	4	4	4	0	0	-	0
"Blue" Wastestream 10 4 4 2 2 0	"Archived"	"Red" Wastestream	10	4	4	0	0	0	0	0	0	0
"Charcoal" Wastestream 10 4 4 0	Wastestreams	"Blue" Wastestream	10	4	4	2	2	0	0	0	0	0
L 5 4 4 4 4 4 6 0 1 3 3 4 HN 5 4 4 4 4 4 0 0 0 0 0 0 0 0 0 0 0 0 0 0		"Charcoal" Wastestream	10	4	4	0	0	0	0	0	0	0
HN 5 4 4 4 4 4 0 0 0 0 0 0 0 0 0 0 0 HD 5 4 4 4 4 4 0 0 0 0 0 0 0 0 0 0 0 0 0 0		1	2	4	4	4	4	4	0	1	3	0
HD 5 4 4 4 4 4 1 0 1 1 1 Charcoal" Wastestream 25 4 12 0 0 0 0 0 0 0 0 0	08/23/96	H	5	4	4	4	4	4	0	0	0	0
"Charcoal" Wastestream 25 4 12 0 0 0 0 0 0 0 0	"Fresh"	Æ	5	4	4	4	4	4	0	-	-	٥
	Wastestream	"Charcoal" Wastestream	25	4	12	0	0	0	0	0	0	0

Note: All times to decontamination were 1 hr.

a Ulceration at some dosing sites may have obscured evidence of microvesication.

b Incidence of sign was significantly less than that for sites dosed with L using McNemar's Test and a significance level of p=0.05.

c Incidence of sign was significantly less than that for sites dosed with HN using McNemar's Test and a significance level of p=0.05.

d Incidence of sign was significantly less than that for sites dosed with HD using McNemar's Test and a significance level of p=0.05.

e Agent (L, HN, HD) at a concentration of 10% in chloroform.

3.3 Data Analysis Results

3.3.1 Gross Pathology (Erythema and Edema)

Means and standard deviations were calculated for erythema and edema scores (Phase II and III Studies) and for lesion areas (Phase III Studies). Analysis of variance was performed for inflammation scores and lesion areas. Table 6 presents means and standard deviations for erythema and edema scores. Significant decreases in average inflammation scores resulted when comparing wastestream-dosed ("archived" or "fresh" -25 μ L volume application) to agent-dosed sites (HD or agent/chloroform) - refer to Table 6. Some significant increases in lesion areas were noted with wastestreams, presumably due to the larger volume dosed. For the "August 13, 1996" experiment (vesicancy assay of "archived" wastestreams), significant decreases in average inflammation scores as well as average lesion areas resulted when comparing wastestream-dosed ("archived" "Red" and "Blue" process wastestreams - $10~\mu$ L volume applications of wastestreams and agent/chloroform solutions) to agent-dosed sites. All observed inflammation scores and lesion areas from the "fresh" "Charcoal" wastestream-dosed sites were zero.

3.3.2 Histopathology

Statistical analysis (McNemar's test) of the histopathology data was performed to ascertain the significance between treatment groups (neat HD, agent/chloroform solutions, and wastestreams) at the 0.05 significance level. Sites dosed with "Red" or "Charcoal" wastestream ("archived", 25 μ L volume application) exhibited a significant decrease in incidence (incidence = 0) of microblisters compared to those sites dosed with HD or agent/chloroform solutions. Sites dosed with the wastestreams also showed a significant decrease in the incidence of follicular necrosis compared to sites dosed with any of the three agents (HD, HN, or L in chloroform; neat HD). Some significant neutralized wastestream versus agent differences also resulted with respect to incidence of epidermal and dermal necrosis.

Sites dosed with "Red" wastestream ("fresh", 25 μ L volume application) showed a significant decrease in incidence of microblisters (incidence = 0), epidermal necrosis, and follicular necrosis compared to that on sites dosed with any of the three agents. Numerical reductions in some pathology from wastestream-dosed sites ("archived", 10 μ L volume application) were observed, although they were not statistically significant due to the smaller number of animals tested.

Statistical analysis of incidence of intermediate to severe histopathologic signs was also performed. Sites dosed with "Red" or "Charcoal" wastestream ("archived", $25~\mu$ L volume application) demonstrated a <u>significant decrease</u> in incidence (incidence = 0) of microblisters compared to that on sites dosed with L/chloroform and HN/chloroform. A decrease in incidence (incidence = 0) was also observed for the "Red" or "Charcoal" wastestream compared to that on sites dosed with HD/chloroform, but were not statistically significant because only four of the

eight animals exposed to HD/chloroform had intermediate to severe microblisters. Sites dosed with "Red" or "Charcoal" wastestream ("archived", $25~\mu L$ volume application) demonstrated a significant decrease in incidence of epidermal necrosis and follicular necrosis compared to that on sites dosed with any of the three agents. Sites dosed with "Red" wastestream ("fresh", $25~\mu L$ volume application) showed a significant decrease in incidence of microblisters (incidence = 0), epidermal necrosis, and follicular necrosis compared to that on sites dosed with any of the three agents. Sites dosed with "Blue" wastestream showed a significant decrease in incidence of follicular necrosis compared to that observed on sites dosed with any of the three agents.

Sites dosed with "fresh" "Charcoal" wastestream (25 μ L volume application) exhibited a numerical reduction in incidence (incidence = 0) of microblisters, although this was not statistically significant due to the smaller number of animals tested, compared to that observed on sites dosed with any of the three agents. Statistical analyses also were conducted on the pooled "Charcoal" wastestream data ("fresh" and "archived", 25 μ L volume applications- see Table 14). These analyses assumed that the probability of a microblister and other histopathologic endpoints is similar for sites dosed with "archived" and "fresh" "Charcoal" wastestreams. Pooled data for sites dosed with "Charcoal" wastestream showed a significant decrease in incidence of microblisters (incidence = 0) and follicular necrosis compared to that on sites dosed with any of the three agents. Statistical analyses of incidence of intermediate to severe histopathologic signs (Table15) were also performed on the pooled "Charcoal" wastestream data ("fresh" or "archived", 25 μ L volume application). Sites dosed with "Charcoal" wastestream showed a significant decrease in incidence (incidence = 0) of intermediate to severe microblisters, epidermal necrosis, and follicular necrosis compared to that observed on sites dosed with any of the three agents.

PHASE III. SUMMARY OF HISTOPATHOLOGY FOLLOWING DOSING OF "CHARCOAL" WASTESTREAM TABLE 14.

	Dose					Number	Number of Animals with Histopathology	pathology		
Agent/ Compound	Volume (µL)	No. Of Animals	No. of Sites	Micro- blister	Epidermal Necrosis	Foilicular Necrosis	Pustular Epidermitis	Dermal Necrosis	Hemorrhape	Vascular
LÉ	5	12	12	12	12	12	0	7	6	
HN	5	12	12	110	12	12	4	. ~	-	
[€] HD	2	12	12	116	12	12			2	
"Charcoal" Wastestream	25	12	20	0 c.d.e	10	4 c.d.e	3	0.0	,0	0

Note: All times to decontamination were I hr.

a Pooled data from the "Charcoal" wastestream received 1/25/96 and dosed on 3/13 and 3/21/96 and wastestream received 8/29/96 and dosed the same day. Volume of "Charcoal" wastestream dosed was

Marked ulceration at the dosing site on animal #496 may have obscured microvesication.

Incidence of pathology was significantly less than that for sites dosed with L based on McNemar's Test at the 0.05 significance level.

d Incidence of pathology was significantly less than that for sites dosed with HN based on McNemar's Test at the 0.05 significance level.

Agent (L, HN, HL) at a concentration of 10% in chloroform.

PHASE III. SUMMARY OF INTERMEDIATE TO SEVERE HISTOPATHOLOGY FOLLOWING DOSING OF "CHARCOAL" WASTESTREAM " TABLE 15.

	Dose	e di s		1 to	Num	ther of Animals wi	Number of Animals with Histopathology Rated Intermediate to Severe	ted Intermediat	e to Severe	
Agent/ Compound	Volume (µL)	No. of Animals	No. of Sites	Miero- blister	Epidermai Necrosis	Follicular Necrosis	Pustulār Epidermitis	Dermal Necrosis	Hemorrhage	Vascular Necrosis
$\Gamma_{ m e}$	5	12	12	10	12	12	0	7	5	0
HNe	5	12	12	10	12	12	0	3	0	0
нD _е	5	12	12	ec	12	12	0	∞	-	0
"Charcoal" Wastestream	25	12	20	0 b,c,d	2 b.c.d	p'o'q O	0	P*40	0	0
Note: All times to decontamination were 1 hr	aminotion were	1 hr								

Pooled data from the "Charcoal" wastestream received 1/25/96 and dosed 3/13 and 3/21/96 and wastestream received 8/29/96 and dosed the same day.

Incidence of pathology was significantly less than that for sites dosed with L based on McNemar's Test at the 0.05 significance level.

Incidence of pathology was significantly less than that for sites dosed with HN based on McNemar's Test at the 0.05 significance level.

Incidence of pathology was significantly less than that for sites dosed with HD based on McNemar's Test at the 0.05 significance level.

Agent (L. HN, HD) at a concentration of 10% in chloroform.

4. Discussion

The intent of the process chemistries was to develop neutralization reactions that achieved destruction of CAIS agents, forming wastestreams with minimal toxic hazards. Achieving the desired objectives represented a formidable challenge since chemical reactions with the agents can result in the formation of reaction products/by-products having vesicant action and/or a high degree of systemic toxicity. Destruction of agents involves complex chemical reactions. The toxicity of the degradation products resulting from the chemical neutralization of HD, HN, or L is of concern to the toxicology, health, and regulatory communities. The current studies were undertaken to assess the vesicant properties of neutralized CAIS.

Current methods for demilitarizing CAIS are still based largely on chemical neutralization via oxidizing materials. The oxidation of sulfur mustard, as pointed out by Franke (1967), represents one of the most important decontamination reactions for HD. The oxidation of sulfur mustard via various oxidizers (e.g., hydrogen peroxide, hypochloric acid and its salts, potassium permanganate, nitric acid, DCDMH, etc.) yields various compounds whose composition depends on the nature of the oxidant used and the specific reaction conditions. Most easily formed is HD sulfoxide which on oxidation yields HD sulfone - both represent major oxidation products of sulfur mustard.

The oxidation of HD not only alters the skin-damaging properties of HD but the systemic toxicity of sulfur mustard as well. The oxidation of HD is of great interest since sulfoxide formation, on chemical neutralization of HD, can be considered a "detoxification". In contrast, the formation of mustard sulfone, a product of further oxidation, can contribute to an enhanced systemic toxicity and vesicant potential of the product solution/mixture. HD sulfone, having the S(O)₂ functional group, is highly poisonous and comparable in toxicity to HD⁴. Research conducted since Philips' review (Philips, 1950) on sulfur mustard pharmacology/toxicology demonstrated that HD sulfone is a highly toxic vesicant.

Certainly, based on the known toxicity characteristics of mustard sulfone, mustard sulfoxide, and their vinyl derivatives; it is crucial that the process chemistries developed for the destruction of CAIS employ oxidants that minimize the formation of HD sulfone and HD analogs having comparable biological activity (systemic toxicity and vesicancy) to that of HD.

⁴ HD is easily destroyed by all chlorinating agents (aqueous or anhydrous medium). Under appropriate conditions, the chlorination of HD can proceed to form various polychlorides. In the presence of water, chlorination of HD is altered resulting in the formation of sulfoxides (Aleksandrov, 1969). Sulfoxides may undergo further oxidation to sulfones.

The vesication potential of HD degradation products/by-products is of concern-information pertaining to sulfur mustard products/by-products is summarized in Tables 16 and 17. The reader is referred to a review on the subject matter (Olajos *et al.* 1996).

Degradation product(s) of nitrogen mustards have not been implicated as having vesicant potential. The principal degradation product of lewisite, namely L oxide, is a potent vesicant.

The vesicant potential of sulfur mustard derivatives (oxidation and chlorination products) has been investigated since the 1920's. Research has indicated that the strongest vesicant action is exerted by β-halogenated sulfides. The position and degree of chlorination influences the vesicant potential of the thioether molecule. With respect to the site of chlorination, Kirner (1928) and Dawson and Wardell (1930) concluded that compounds having the chlorine atom in the beta position were considerably more vesicant that those having chlorine in the alpha or gamma position. The degree of chlorination also influences the vesicant activity of the sulfide molecule and hence the early use of chlorination to degrade HD. Monosubstitution analogs of HD. regardless of position, are less effective vesicants than HD. As previously stated, the introduction of halogen atoms results in decreased toxicity and markedly diminished vesicant action. Research in the 1920s summarized by Bouder (1940) - indicated that the higher chlorinated derivatives (e.g., tri-, tetra-, and hexachloro derivatives) of HD (saturated or unsaturated) were non-vesicant. A summary of the vesicant potential of various chlorinated analogs of sulfur mustard are given in Table 17. Fuson et al. (1943) on review of the vesicant activity of sulfur compounds concluded that compounds containing the S(0) group were nonvesicant. Mustard sulfone, containing the S (0)2, functional group is a known vesicant (vesicancy potential 1/7 to 1/5 of HD; Bergmann et al., 1945). The formation of HD sulfone can contribute to an enhanced vesicant potential of the product solution/mixture (wastestream).

The lack of vesicancy following treatment with "Red" and "Charcoal" process wastestreams is indicative of the effectiveness of the neutralization chemistries in destruction of chemical agent concomitant with the minimization of potentially vesicant-inducing products/by-products. The composite agent (HD, HN and L) levels in "archived" and "fresh" "Red" wastestreams and in "archived" and "fresh" "Charcoal" wastestreams did not elicit vesication in the volumes dosed. Treatment with "Blue" process wastestreams ("archived" and "fresh") resulted in a vesicant response. The bioassay results were unexpected since the agent (HD) residual level was below 50 ppm, a level not expected to elicit a vesicant response. The most plausible explanation is the presence of vesicating product(s)/byproduct(s).

SYNOPSIS OF DERMAL TOXICITY DATA FOR CAIS AGENTS, AGENT DEGRADATION PRODUCTS, RRS OXIDANT, AND SOLVENTS* TABLE 16.

Compound	Dermal Toxicity ^b (LD _w /LDLo/TDLo)	References	Skin Effects (Irritation, Vesication)	References
AGENTS HD [bis(2-chloroethyl)sulfide]	LD ₅₀ (40-100 mg/kg)	Anslow & Houck (1946)	Severe irritant/escharotic, severe vesicant	Marshall & Williams (1921); Gates & Moore (1946);
L [dichloro(2-chlorovinyl)arsine]	$\mathrm{LD}_{\mathrm{Su}}$ (5-6 mg/kg)	Cameron et al. (1946); Gates et al. (1946)	Severe irritant/escharotic, severe vesicant	Kensnaw (1940) Gates et al. (1946)
HN-1 [bis(2-chloroethyl)ethylamine]	LD ₅₀ (15-20 mg/kg)	Smith (1943a); Anslow & Houck (1946)	Severe irritant/escharotic, severe vesicant	Cope <i>et al.</i> (1946); Renshaw (1946)
HN-3 [tris(2-chloroethyl)amine]	LD ₅₀ (5-20 mg/kg)	Smith (1943d); Anslow & Houck (1946)	Severe irritant/escharotic, severe vesicant	Cope <i>et al.</i> (1946); Renshaw (1946);
OXIDIZED DERIVATIVES HD sulfoxide	· •••	(-)	Irritant, non-vesicant	Marshall & Williams (1921); Lawson & Dawson (1927); Young et al. (1944)
Sulfoxide, 2-chloroethyl vinyl	P (-)	5 0	Irritant, non-vesicant	Thomson et al. (1945)
Divinyl sulfoxide	•	(-)¢	Irritant, non-vesicant	Fuson et al. (1943); Young et al. (1944); Thomson et al. (1945)
HD sulfone	₹	f -)	Irritant/escharotic, vesicant	Marshall & Williams 1921); Young et al. (1944)
Sulfone, 2-chloroethyl vinyl	•	* -	Irritant/escharotic, vesicant	Young et al. (1944); Thomson et al. (1945)
Divinyl sulfone	LD ₅₀ (* 20 mg/kg)	Smyth et al. (1962)	Irritant/escharotic, vesicant	Young et al. (1944); Thomson et al. (1945)
HN-1 oxide	4 (-)	4 O	" (-)	u(÷)

TABLE 16. (Continued)

Compound	Dermal Toxicity* (LD ₅₀ /LDLo/TDLo)	References	Skin Effects (Irritation, Vesication)	References
OXIDIZED DERIVATIVES (Cont.)	7			
HN-3 oxide	! (-)	·(•)	u(-)	"(·)
Lewisite oxide	, (-)	(-)	Irritant/escharotic, vesicant	Young et al. (1944); Thomson et al. (1945)
2-chlorovinylarsonic acid	* (-)	* (-)	Irritant, non-vesicant	Young <i>et al.</i> (1944); Thomson et al (1945)
2-chlorovinylarsonous acid	<u>-</u> C	·-(-)	Irritant, non-vesicant	Cameron <i>et al.</i> (1946)
OXIDIZERS				
DCDMH	LD ₅₀ (>20 g/kg)	EPA 8E-8100-228 EPA 88-8100-228	Severe irritant	EPA 8EHQ0281-0382; EPA #88-8100-173 (cited in RTECS)
SOLVENTS				
Chloroform	LD _{s0} (>20 g/kg)	NTIS AD-A062-138	Mild irritant (cited in RTECS)	Guido and Martins (1988)
t-butyl alcohol	w (-)	u(-)	Mild irritant	Oettel (1936)

Table modified from that originally compiled by Olajos et al., 1996.
 Rabbit as animal model unless otherwise indicated. Tests for irritancy based on animal and/or human studies.

Test for vesicant action of agents conducted on human subjects.

Mouse s.c. LD₅₀ (>25 mg/kg) [Anslow and Houck (1946)].

Rat oral (100 mg/kg, mortality 1/1) [Young *et al.*, 1944]

Mouse s.c. LD₅₀ (>25 mg/kg) [Anslow and Houck (1946)].

Mouse s.c. LD₅₀ (>25 mg/kg) [Anslow and Houck (1946)].

Acute toxicity undetermined.

Mouse i.p. LD₃₀ (50-100 mg/kg) [Bergmann and Fruton (1943); Stahmann and Bergmann (1946a)]. Mouse i.p. LD₃₀ (2-5 mg/kg) [Bergmann and Fruton (1943); Stahmann and Bergmann (1946a)].

Mouse s.c. [mortalities: 2 mg/kg (0/5); 5 mg/kg (5/5); 10 mg/kg (5/5)] Young et al. (1944), Mouse i.p. [mortalities: (1000 mg/kg 10/10; 500 mg/kg 0/10] (Young et al., 1944). Reported as highly toxic, details not given (Cameron et al., 1946).

"Rabbit oral LDLo (4.5 g/kg) [RTECS].

[&]quot; Young et al., (1944) reported HN2 oxide as non-vesicant; no data for HN1, HN3.

TABLE 17. VESICATION POTENTIAL OF VARIOUS ANALOGS/ DERIVATIVES OF SULFUR MUSTARD^a

Analogs/Derivatives (Saturated and Unsaturated)	Vesicant Activity	References ^b
OXIDIZED DERIVATIVES		
Mustard Sulfone (sulfone, bis(2-chloroethyl)	(POS)	Marshall & Williams (1921), Young <i>et al.</i> (1944)
Sulfone, 2-chloroethyl vinyl	(POS)	Young et al. (1944)
Divinyl Sulfone	(POS)	Young et al. (1944), Thomson et al. (1945)
Mustard Sulfoxide (sulfoxide, bis(2-chloroethyl)	(NEG)	Marshall & Williams (1921) Lawson & Dawson (1927) Fuson et al. (1943) Bergmann et al. (1945)
Divinyl Sulfoxide	(NEG)	Young et al. (1944) Thompson et al. (1945) Bergmann et al. (1945)
β-chloroethyl vinyl sulfoxide	(NEG)	Young et al. (1944)
α,β' -trichlorodiethyl sulfoxide	(NEG)	Young et al. (1944)
CHLORINATED DERIVATIVES		
bis(α-chloroethyl) sulfide	(NEG)	Peters and Walker 1923) Baldwin et al. (1924) Kirner (1928) Dawson & Wardell (1930)
α , β , β' -trichlorodiethyl sulfide	(NEG)	Mann & Pope (1922) Lawson & Dawson (1927)
$\alpha,\beta,\beta,\beta'$ tetrachlorodiethyl sulfide	(NEG)	Mann & Pope (1922) Lawson & Dawson (1927)
α , α' , β , β' tetrachlorodiethyl sulfide	(NEG)	Lawson & Dawson (1927)
$\alpha,\alpha\beta,\beta,\beta'$ hexachlorodiethyl sulfide	(NEG)	Mann & Pope (1922) Lawson & Dawson (1926) Dawson & Wardell (1930)
$\beta\text{-chloroethyl}\ \alpha,\beta$ dichlorovinyl sulfide	(NEG)	Lawson & Dawson (1926) Kirner (1928) Dawson & Wardell (1930)
β -chloroethyl α , β , β' trichlorovinyl sulfide	(NEG)	Lawson & Dawson (1926) Kirner (1928) Dawson & Wardell (1930)
β -chloroethyl chlorovinyl sulfide (α and β isomers)	(POS)	Lawson & Dawson (1926) Dawson & Wardell (1930) Fuson et al. (1943)

a Table from Olajos et al., 1996

 $^{^{\}rm b}$ Citations are primary and/or secondary

5. Conclusions

Based on the findings of these studies the following conclusions can be made.

- The vesicating properties of the "Blue" wastestream (product solution from neutralized neat HD) were not significantly reduced from that of the untreated CAIS (neat HD) prior to treatment with neutralization solution.
- The vesicating properties of both "Red" and "Charcoal" wastestreams (product solutions from neutralized agent/CHCl₃ and agent/charcoal, respectively), in the volumes dosed, were significantly lower than the untreated CAIS agent solutions.
- The microvesicancy test results on the "archived" wastestreams and "fresh" wastestreams suggest that storage had not altered the vesicancy potential of the product solutions (wastestreams).

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APPENDIX A

Gross Lesion Appearance (24-hr)

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Project #: <u>G1555</u> -	-38A					Date:	2-20-6	16
MREF Protocol #:	109				Study Di	rector: Carl Oli	500	
Day: 2		Lesion	Read By:	75	_ Lesions	Recorded By: _	Rmm	
Lesion Sites	A	С	Ε	G	В	D	F	COMMENTS
Animal I.D. #								
301	15/10	150	16/14	13/14	22/23	19/32	15,0 R-3 E-3	readings Takin in mm
	E432 R-2	K-22	E-3 R-3	R-2 E-2	R-3 E-2	R-2 2-2	R-3 E-3	
								·
Mean Average								
All Measurements N/A = Nct applies N/It = Not require	able	rs.	R = Erythe E = Edems 1 = Mild 2 = Mode 3 = Severe	rate	OW.	2-20-9	6 EMM	
Stie A 10ml	1070HD	<u>CHC</u>	3.					
Site B 50.2	102 HD	in CHC	Z =					
Site C_10.0	10% HV.	in CHC	£	•				
Site D 50.2	14 JOI	Sim CHO	43					
Site E	107-1	Lin CH	<u>L</u>					
Site F 50.0	ich L	in CH						
Site G \ L	H tain	<u>(T</u>						•
Reviewed By:	77	DP 32		Date:	2/2	3/7:		·
Anne	ndix A			55	5	•		

Appendix A

Project #: <u>G1555</u> -	-38A	·				Date:	2-20-9	<u> </u>
MREF Protocol #:	109				Study Dir	ector: <u>Carl Ols</u>	son	
Day: 2		Lesion	Read By: C	<i>3</i> 0	_ Lesions R	lecorded By: _	pm.nr	
Lesion Sites	A	С	Е	G	В	D	F	COMMENTS
Animal I.D. #								
305	108	12/2	9/16	14/3	19/20	23	31/25	readings taken
	R-2 E-2	R-3 €-3	R-2 E2	R-2 E-2	R-3	R-3 =-3	683	
Mean Average								
All Mearttrements in N/A = Not applical N/R = Not required	ole	5.	R = Erythe E = Edena 1 = Mild 2 = Moder 3 = Severe	-ate				
Stie A 10-0 10								·
ike B <u>50 nd 1</u>								
lite C 10 m 1								
ite D_50~0 \			_					
ire = 10.0_								
ite F_50.L			HCL3					
ite G <u>lad</u>	then	HD					••	
eviewed By:	176	<u> </u>		Date:	2/2	3/3.		
Appen	dix A			56				

Project #: <u>G1555</u> -	38A					Date:	7-22-	<u>96</u>
MREF Protocol #:	109				Study Di	rector: Carl Ols	ion	
		I esion	Read By:	S. K	Lesions F	Recorded By: _	iamm	
Day: 2					<u> </u>			GOV 0 / EVITS
Lesion Sites	A	С	E	G	В	D	F	COMMENTS
Animal I.D. #					<u> </u>			1,500 \
306	15/8	78	78	16/7	16/3	12-14	12/0	in mon
J () e1	R-22 E-3	R-3 E-3	R·1 E-2	R-2 E-3	R-2 E3	R=32 E-1	R-2 E-3	
	2-0			1				
				·				
·		·						
Mean Average								
All Measurements N/A = Not applim N/R = Not require	b!	s.	R = Eryth E = Edem: 1 = Mild 2 = Mode	rate			0IF 1-	22-96 Amm
Stie A 5.0.10	270 L in	- CHCla	3 = Severe					
Site B Cas 10						•		
Site C 5-0 10	To HD	incho	<u>-1</u> 3					
Site D Cond 1								
Site E _ 5-2-1								
Site F 1 Cal			<u>/3</u>					
Site C 1 no	H true	<u>D</u>						
	J. L	(M.,			2/23	176		
Apper	ndix A			57				•

Project #: G1555	-38A					Date:	2-2.2-5	<u>ic</u>
MREF Protocol #				4	=	ector: Carl Ols		
Day: 2		Lesion	Read By:	10E	_ Lesions R	lecorded By: _	BMM	
Lesion Sites	A	С	E	G	В	D	F	COMMENTS
Animal I.D. #								
309	16/0	12/3	15/14	1410	79	99	7/0	readings mm
	R-22	R-1 E-2	R-3 E-3	R-1 E-2	R-3	R-22	RE-SIM	
·								
		÷						
	:				-			
Mean Average								
All Measurements N/A = Not applies N/K = Not equire	ble	<u>.</u>	R = Erythe E = Edem: 1 = Mild 2 = Mode 3 = Severe	nate	<u>. </u>			
Stie A 10 nd								
Site B 5.02 10								
Site C 1020								
Site D 5-2	1070 H1	Vinct	<u>1</u> 43					
Site E !OLC								
Site F 5 2 2			<u> </u>					
Site C Luce	nent	HD						
Reviewed By:	(1. 1	[1]-		Date:	2/23/	34		

Project #: <u>G1555</u> -	-38A					Date:	2-28-	96
MREF Protocol #:	109				Study Dir	ector: Carl Ol	son	
Day: 2			Read By:	7.	_ Lesions R	lecorded By: _	N-MCKIL	
Lesion Sites	A	С	E	G	В	D	F	COMMENTS
Animal I.D. #								
312	15/12	1910	14/2	2015	9/2	9/11	123	frederings token in mm
212	R-3	£-2 €-2	R-3	R-3	R-3	R-3	K. 12 2	
Mean Average								
All Measurements N/A = Not applica N/R = Not require	ible ed.		R = Erythe E = Edenia 1 = Mild 2 = Moder 3 = Severe			00001	0-23-96	omm
Stie A / Cul /	0% L.	in CHC	13					
Site B <u>5 ul / l</u>	0% L 1	(H(1	<u>3</u>					
Site C_10.11 / 6								
Site D <u>5ul 10</u>								
Site E 10 ml 10	70 HN	<u> </u>	<u>C1</u> 3					
Site F 5 40 /0			<u>C/3</u>					
Site G/W								
Reviewed By.		<u>()15</u>			2/25/	٤٤		·
Apper	ndix A			59				

Project #: <u>G1555-</u>	38A					Date:	2-25-9	6_
MREF Protocol #:					Study Dir	ector: <u>Carl Ols</u>	on	-
Day: _2	• • •	Lesion	Read By:	170	_ Lesions R	ecorded By:	insince_	
Lesion Sites	Α	С	Е	G	В	D	F	COMMENTS
Animal I.D. #								
316	13-9	15/3	31/4	3214	11/9	15/9	14/14	Kindengo taken im mm
	R-2 E-2	8-3 8-3	R-3	R-3	R2 =3	R-3 g 3	R-3 E-2	
Mean Average								
All Measurements in N/A = Not applicate N/R = Not required	ole i.		R = Erythe E = Edema 1 = Mild 2 = Moder 3 = Severe	rate	·			
Stie A 1022 Stee B FILL 1								
Site B <u> </u>								
Site D <u>Ful /C</u>								
Site E 1040/14								
Site F 5216/16								
Site G / 2: ()								
Reviewed By:	(T	Ofin		Date:	2/23	156		

		LESIC	ON SIZE	DETFI	RMINA.	110N 21	TEET			
Project #: <u>G15</u>	Ф _{ЗВА} 55 -9001	}					Date:	_3-6.	-96	
MREF Protoco	1#: 109)		_ Stud	y Direc	tor: <u>Car</u>	l Olson			
Day:	L	esion Re	ead By:	(,0	I	esions R	Recorde	d By:	JMIH	_
Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS]
Animal I.D. #					-	<u> </u>		1.14	C: a dian	-
313	1013	100	12/14	14/16	NIANIA	NANIA	NANA	NINA	ELEMENT	1
	R- 2	R-2 E-3	R-3 E-3	F-3		E-1	E-0'	E-1		1
	15-1				P-0	HINCES				
· · · · · · · · · · · · · · · · · · ·										
	-									1
	 									1
										<u>.</u>
						-				1
										-
									2.10.00	<u> </u>
All measurements in N/A = Not application N/R = Not require	able	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe	a 0	= Notay AC N	pparest NA in n-thi nct t	t egs in for	ivalent in throw tidy 10-2	to 0 1gh 14-96 01
Site A <u>5 ul 100</u>	z HD	in Ct	(Cl3						đ	
Site B <u>20ul n</u>		_		5n						
Site C <u>5ul 10</u>										
Site D <u>20uln</u>				, m						
Site E <u>5 ul 1</u> (
Site F 20 il M				-						
Site G / ul s										
Site H <u>2Oul</u> n			, solii	tion						
Form No. MREF-LES		_					\sim	1 1	17 Din	
OEnstyérias	5-6-176.	المنط					انعاديه	معد صر 11ء	و آ کائے ما	, _
<u> </u>	2				63			•		

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OENGLYELEN, 5-6-170 SAIN BEE 3-6-46 SANH Appendix A

Project #: <u>G15</u>	Ф _{ЗВ.4} 55- 9001	1	JN 3121	E DE LEI	KMINA	110N SF		3-6	-96	
MREF Protocol	#: <u>109</u>)		_ Stud	ly Direct	or: <u>Car</u>	l Olson			
Day:2_	L	esion Re	ad By:	وش	L	esions R	ecorded	i By: 🗻	Incit	_
Lesion Sites	A	С	E	G	В	D	F	н	COMMENTS	
Animal I.D. #			,	,	6	2)	1 12 2			
315	150	12 13	11 9 R- 2	18 12	NA NIA	NIA E-O/A	NIA NA	NANA R-O	talement mm	
	R-1 F-1	L-3 E-3	E-3	足-み E-3	E-1	E-1	E-1	E-1		
							1			
				·					<u> </u>	
				-					•	
All measurements in N/A = Not applicate N/R = Not required	ole	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe		0	= Y) + AC ,	DALO WHILL The	rent OWN & equivale from the	nt to to a rough of AGA
Site A 5 ul 10 %	3 HD.	mCH(213							
Site Balulma	utral	·	olutio	7						
Site C 52l 10%		<u> </u>								
Site D 20 wl Me				ion						
Site E <u>5ul 107</u>	o HD.	nCHC	.13							
Site F 2000 me				tion						
Site G/ul ne	at H.	<u> </u>								•
Site H <u>20ulme</u>	utial	13119	roluti	ial.						

Form No. MREF-LESION.SIZ-07

Project #: <u>G155</u>	0 5- 9001						Date:	3-6	.96
MREF Protocol	#: 109	1		_ Stud	y Direct	or: <u>Car</u>	l Olson		
Day:			ad By:	CY	<u>o</u> L	esions R	ecordeo.	i By:	<u>lmit</u>
Lesion Sites	Α	С	E	G	В	D	F	Н	COMMENTS
Animal I.D. #							1 - 4		100 dina
317	9/15	8 8	13/12	15/10	20	20	20	00	Taken Mu
·	R-3 E-3	トコ	L- 3 E- 7	L-3 E-2	R-0.	R-0 E-1	R=0 =-1	E-0 E-1	
·									
			·						
						•			
					·				
							<u> </u>		
All measurements in N/A = Not applicat N/R = Not required	ole	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe	. O=	Mot C	Ippne	ut @	WN 3-6-96 BMM TE 3-6-96 BMM
Site A 511/07	Lin	CHCIS	,						
Site B 20ul ne		_		L					
Site C <u>5ul 107</u> 0									
Site D 20 ul no	utial	inging!	poletin	u.					
Site E <u>5 wl/07</u> 0	<u>HVin</u>	CHC	13						
Site F Zilul Me	utra	lizing	soluti	را. ماوسی					
Site G / wl ne	at t	HD_							•
Site H <u>20 ul 1</u>	west.		تنتضمر.	teon					

Form No. MREF-LESION.SIZ-07

Project #: G15	0 ₃₅ q 55- 9001		JN 3121	י ארי בי	MINA I	IION SF		2-6	·- 96
MREF Protocol	#: 109)		_ Stud	y Direct	or: <u>Car</u>	l Olson		
Day:	L	esion Re	ead By:	Cro	L	esions R	ecordeo	i By:	<u>Jm}</u>
Lesion Sites	А	С	E	G	В	D	F	Н	COMMENTS
Animal I.D. #						,			
324	9/12	10/1	13 14	149	00	00	25	00	tike imm
	£-3 E-3	2-2 E-2	R-3 E-3	R-3 E-3	R-0 E-0	E-1	R-0 E-1	R-0 E-1	
					·				
			,						
All measurements in N/A = Not applicat N/R = Not required	ole		E = 1 = 2 = 3 = 3	Erythema Edema Mild Moderate Severe	Ð=	Not ap	pricat		WN 3-6-96 DRIN
Site A <u>5 ul 10 7</u> 0	HDi	n CHC	1/3						
Site B 20 rd ne	utral	izing D	polution	-					
Site C 5 21 1070	HNin	<u>CHCl3</u>							
Site D 20 ul ne	utral	izing D	olution						
Site E 510 10%	Lin C	HC13							
Site F. 20. nl nee	itrali	zing s	olution	_					
Site G / Ll Men								•	

Site H 20 ul noutralizing solution Form No. MREF-LESION.SIZ-07

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Revenued by CT Off

Project #: G15	55- 900 	0					Date:	3-6	6-96	
MREF Protocol	#: 109)		_ Stud	ly Direct	or: <u>Car</u>	l Olson			
Day:	L	esion Re	ad By:	<u>CTO</u>	L	esions R	Recorded	d By:	Inch_	
Lesion Sites	А	С	E	G	В	D	F	н	COMMENTS	ı
Animal I.D. #										
3//	13/1	149	13-9	15/10	00	00	20	200	tedings	
	2-2	R-3 E-3	R-3 E-2	R-2 E-2	R-0 E-1	₽-0 E-1	1-0 E-1	R-0 E-1		
									•	
All measurements in N/A = Not applicat N/R = Not required	ole		E = 1 = 1	Erythema Edema Mild Moderate Severe	0:	e Net	арраі	ent E	שאל 3-6-96 kg	PALICE FALAC
Site A <u>5ul 107</u> 5										
Site B <u>20ul Meu</u>	itrali	zing si	lution							
Site C <u>Jul 1070</u>										
Site D <u>. 20 ul me</u> u		•		ıL						
Site E <u>5_1076</u>	H Din	CHC/3								
Site F <u>20 ul No</u> Site G <u>Dul Ne</u>	eutra et Hi	lizing D	soluti	in						
Site H <u>2010 m</u>			معنى نين	.						

Form No. MREF-LESION.SIZ-07

Project #: G15	55- 38 <i>8</i>	A				Date:	3-	14-96	
MREF Protoco	1#: <u>10</u>	9 P.h	e 3	Stud	dy Direct	tor: <u>Car</u>	ri Olson		
Day:2_	I	Lesion R	ead By:	B	L	esions F	Recorded	d By: <u>LOM</u>	u.
Lesion Sites	A	С	E	G	В	D	F	COMMENTS	
Animal I.D. #	0								
310	\$ 13	98	12/0	910	1722	11/17	15/6	seedings taken in mm	
	R-1 E-2	R-1 E-2	R-3 E-3	R-3	R-1 E-2	R-1 E-2	R-05-Q	·	
·									
·	<u> </u>								
	<u> </u>								
			0=	not an	MASSINE	OFF	3-14-96	· Emm	
All measurements in		ers	R =	Erythema		@AC_	The l	esion rea	dingo Were Liveld duae dae 0.5. 90 our
N/A = Not applicat N/R = Not required			1 = 1	Edema Mild		letu	ven Ø.	mined at 0 and iar	dwas_
•			2 = 1	Moderate Severe	٠.	there	fore.	designate	der C.J.
- 0 1007	ИD.	с цс	7 1-			BEE 1	0-16-96	Rum	•
Site A <u>5 ul /07</u>									
Site B 25 ul R	لعالم	avlast	ream						
Site C <u>Jul 107</u> z	HNin	-CHCI	3						
Site D <u>25ulB</u> s	en W	aster	tream						
Site E <u>5ul/0%</u>	Lin(CHC13							
Site F <u>25ulCh</u>	arra	l War	testre	<i>م</i> ،-					
Site G/ul ne	atH.	<u>D</u>							
							a .	,	\mathcal{L}^{ι}

Form No. MREF-LESION.SIZ-07

Appendix A

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Project #: G15	55- 38A					Date:	3-/	14-96	
MREF Protocol	#: 109	Pha	<u>u3</u>	_ Stud	ly Direct	or: <u>Car</u>	l Olson		
Day:	L	esion Re	ad By:	<u>OB</u>	L	esions F	Recorded	i By: <u>iQM</u> M	<u></u>
Lesion Sites	Α	С	E	G	В	D	F	COMMENTS	
Animal I.D. #			0 /	1.0	10/	<u> </u>	10/	readings	
491	129	11/15	8 10	12/19	805-64	13/7		takurimm	
	R-3 G-3	R-3 G-2	<u>e</u>	R-1 E-3	R 05-6	E-2	R.21 E-2		
	· ·								
	<u> </u>								
<u> </u>									. •
All measurements in N/A = Not applica	ble I		R = E = 1 = 2 =	Erythems Edema Mild Moderate	3)SE 3-	14-96	Dum	a fiece of attacked ald be due our night blass were this
Site A <u>5ul /07</u>				thin p	ule d	Back	2.14-96	, oun co	uld be due
Site B <u>25ulCA</u>			estria	to	tam	~ trai	ma	induded	las were
Site C <u>5 ul /07</u>	, HDin	<u> CHCI</u>		_	y thin	anin	ine a	nimeli o 91. Drim	n this
Site D <u>25ul R</u>	edli	astes	tream						
Site E <u>5 ul 107</u>	o HNi	<u>_CHC</u>	3	ΘA	C Ih	u lea	con n	+ Davido	hetween
Site F 25 ul B				,	. 0,0	مهمه (Llan	dwar th	erefore DNA
Site G <u>/.u.l</u> M					d	esign	atid	مین ن.ع. ع	. ,

Form No. MREF-LESION.SIZ-07

Project #: G15	55- 38A					Date:	3-14	-96	
MREF Protocol	#: <u>109</u>	Pha	se 3	_ Stud	ly Direct	or: <u>Car</u>	l Olson		
Day:2_	L	esion Re	ad By:	B	L	esions R	lecorded	By: _ADMA	<u>u</u>
Lesion Sites	A	С	E	G	В	D	F	COMMENTS	
Animal I.D. #							1		
493	9/7	1/1	19/9	14/15	1500	1919	15/20	Readings taken in MM	
	K-3	6-1	R-3 E-3	R-3 E-3	R-1 E·i	R·1 E-2	R 0,50		
						•			
									4+
All measurements in N/A = Not applicat N/R = Not required	ole .	ers	1 = 2 =	Erythema Edema Mild Moderate Severe	AC AC C	The le Lotern),0 and lesign	sioni uned d I an vated	st livels at livels d was t	were best between herefore 3-14/96 Du.

Site A <u>5ul 1070 H Din CHC13</u>

Site B <u>25ul Red Wastestream</u>

Site C <u>5ul 1070 H Win CHC13</u>

Site D <u>25ul Blue Wastestream</u>

Site E <u>5ul 1070 Lin CHC13</u>

Site F <u>25ul Charcal Wastestream</u>

Site G <u>1ul neat HD</u>

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Appendix A

Revenued by CT Elson 3/18/50

Project #: <u>G1555- 38A</u>					Date: 3-14-96						
MREF Protocol #: 109 Phase 3 Study Director: Carl Olson											
Day:2	L	esion Re	ad By:	B	L	esions R	tecorded	i Ву: _ <u>С</u> м.	<u>u</u>		
Lesion Sites	Α	С	E	G	В	D	F	COMMENTS			
Animal I.D. #									-		
498	79	12/1	910	17/5	1733	RW- 0		Asadrago takinin mm			
	R-3 E-1	K-3	P-3 L3	R-2 C-3	R-3 Ii	EC2	E-3		1		
				. ·	-						
All measurements in millimeters N/A = Not applicable N/R = Not required 1 = 2 = 3 =				Erythema Edema Mild Moderate Severe	TOAC	Jhe diter 0.0 a desig	lesce mine not o notes	n reading eat level and was a as 0.5	Jetu Theref 3 14-46	best he Dav	
Site A <u>Sul 1070 HN in CHC13</u>											
Site B 25ul Blu											
Site C 5 2 109											
Site D 25ul Ch	arcoal	4)ast	istra	200							
Site E 5.4107											
Site F25ul Red Wastestran											
Site G Lul Me	st H	<u>D</u>									

Form No. MREF-LESION.SIZ-07

Revenued by CT Dim 311519x

Project #:	<u>G1555- 38A</u>
------------	-------------------

Date: 3-22-96

MREF Protocol #:	109 Phone 3	Study Director:	Carl Olson
		_	

Day: 2 Lesion Read By: 18 Lesions Recorded By: 10 MM

Lesion Sites	А	С	E	G	В	D	F	COMMENTS
Animal I.D. #						T		,
499	78	1010		12/6	13/5	15/E	9/6	
	R-2 E-2	R-2 E-3	R-3 E-1	Q-3 €-2	R- 1	R-0.5	R-1	
		U=40	4=6	U= 4				
								•
		·			,			
·								
			·					

All measurements in millimeters

N/A = Not applicable

N/R = Not required

= not apparent (AC The lesion reading were best

E = Edema

1 = Mild

2 = Moderate

3 = Severe

O.O and I and was therefore disignated as 0.5. 3-2296 to DAC relevation of dose sites were not perviously record 3-28-96 Emm

determinded at levels between

U= ulceration noted as:

4= small

5= medium

6= large

Site B 25 ul Blue Wastestream

Site C 5 ul 10% Lin CHC/3

Site D 25 ul Charcoal Wastesm

Site E 5 ul 10% H Din CHC/3

Site F 25 ul Red Wastestream

Site A 5 ul 107 HNin CHC/3

Site G/ul weat HO

Form No. MREF-LESTON.SIZ-07 Appendix A

Revenued by CT Elem 3/28/90

						_	000	n 67		
Project #: G15		_						2-96		
MREF Protocol										
Day:2_	L	esion Re	ad By:	R	L	esions R	ecorded	By: <u>iOn</u>	. <u>m</u>	
Lesion Sites	А	С	E	G	В	D	F	COMMENTS		
Animal I.D. #							1 19 1			
494	14/3	1013	10/1	17/5	00	219	1320			
	R-3 E-3	R-2 E-2		R-3 6-3	R-0	R-1	R-1			
			U=4	U=4						
	·								• •	
			•							
					•					
All measurements in N/A = Not applical N/R = Not required	ble	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe		O=N OAC. Not	ulce pre	eparent stime of coursely 3	lose site recorded -28-96 om retec an);	<u>)</u> wi
Site A <u>5ul 10</u>	70 Li	mCHO	2/3					4 = smal 5 = medi	um	
Site B 25 ul Ch				ion				6 = large		
Site C <u>5ul 10</u>										
Site D 25 ul 18				J						
Site E .5 11 10°										
Site F 25 ul B			itrear	w_						
Site G / ul M	eat	40								

form No. MREF-LESION.SIZ-07 Appendix A Personed by CT Dem

Project #: G15	<u>55- 38</u> 4	7				Date:	3-2	2-96			
MREF Protocol	#: <u>10</u>	2 Pho	رون ع	_ Stud	ly Direc	tor: <u>Ca</u>	rl Olson		<u></u>		
Day:2	I	esion R	ead By:	<u>Ac</u>	I	esions F	Recorded	i By: 10 min	<u>~</u>		
Lesion Sites	А	С	E	G	В	D	F	COMMENTS			
Animal I.D. #			·								
496	12/5	109	13/6	17/6	1921	1919	26				
	R-3 €-3	R-3 E-i	R-3 5-3	R-3	R-1 E-2	R-3 E-2	R-0				
	u=60	4=6	4=6	W=4							
·											
				•							
	1								• •		
·					•						
All measurements in N/A = Not applicable N/R = Not required		ers	E = 1 = 2 = 3	Erythema Edema Mild Moderate Severe	in site mot previously rate recinded 3-28-96 6						
Site A <u>5ul 107</u>	L HD	n CHC	:/3			=ىلا:	مەلىر	ration n H=A	sted as:		
Site B 25 ul Rs	ed Wa	estest	eim					5= N	nedium		
Site C 5 ml 10°									arge		
Site D 25 ul Bl	nelva	ratest	ream								
Site E <u>5 2 2 1 0 7</u>	oL in	CHC	/3								
Site F 25 21 (-	harc	<u>oal li</u>)astis	bream			•				
Site G 1 rul n	eath	10									

Form No. MREF-LESION.SIZ-07

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Reserved by Ci Olsen

· .	٠						200	2.01			
Project #: G15	55- 38A	•				Date:	2-2-	2-96_			
MREF Protocol											
Day:2	L	esion Re	ad By:	_9B	L	esions R	ecorded	I Ву: <u>ДМ</u> /	<u>u</u>		
Lesion Sites	A	С	E	G	В	D	F	COMMENTS			
Animal I.D. #			,				1 11 -				
497	8 8		10/12 R-3	15 21 R-3	2120 R:2	R-0.5 6 E-0.5 6	R- 2				
	R-3	R-3 G-3		R-3 G-3	G-2	E-C'2	6-2				
	U=66		n=18	11-6							
			· ·								
									·		
									OIE 3-22-46 EM		
	All measurements in millimeters $R = Erythema$										
Site A <u>5 ul /0°</u>					(3)	ر HC م بد	ulcer	not pri	stose sites viously record.		
Site B 25 ml B	luele) acted	tream				. 3-2	5-7 <i>6 R4 1</i> 1CN	C V		
Site C <u>5 ul 10 1</u>						-		ation no	ted av:		
Site D <u>25 ul Ck</u>	arcoa	l Was	testria			,	= sm	all Lum			
Site E <u>5 rd</u> 109	TO HD	in CH	1013				= lan				
Site F 25 - u & R	red h) astest	rean.								
Site G / <u>/ / / / / / / / / / / / / / / / / </u>	eat.	HD_									

Form No. MREF-LESION.SIZ-07

Project #: G15	Project #: G1555-38A Date: 6-21-96										
MREF Protoco	1#: <u>10</u> 9	9 Pho	<u>E</u>	Stud	dy Direc	tor: <u>Ca</u>	rl Olson				
Day: 2	I	esion R	ead By:	B12-	I	esions I	Recorded	d By:	D. p.Cm		
Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS		
Animal I.D. #											
346	28/2	12/8	109	11 R-2	R-0	1621	00	19 20			
	R-3 E-3	R-2 E-2	5-7	E-2	€0	R-1 E-1	E-0	₽-1			
									·		
									•		
				•							
All measurements in N/A = Not applical N/R = Not required	ble		E = 1 = 2 =	Erythema Edema Mild Moderate Severe		0	= M. 2.	tapp	prest		
Site A <u>5.1107</u> 0	Linc	4013									
Site B 25-4	1	Toutra	-				•				
Site C Sal 1010	~ (TH	<u>~CHC</u>	3								
ite D 25.1 60	ua mo	Jeste									
Site E 5-0 (070	tho in	<u>- CHC</u> I	3								
ine F 25-0 -s	Luci	taites									
ite G <u>lubas</u>	± 47				٠				•		
ite H_25_0 b	<u> </u>	- steet	~~~								
	N 617.07						R	ر ند. حکا	. by		

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Appendix A

U7 QD2 6/24/91

Project #: G15	55- 38A	.				Date:	6-3	21-96	_	
MREF Protocol	#: 109	Pho	<u> 8</u>	_ Stud	ly Direct	or: <u>Car</u>	l Olson			
Day:	L	esion Re	ad By:	BIL	<u> </u>	esions R	ecorded.	By:	Cmm	
Lesion Sites	A	С	E	G	В	D	F	H	COMMENTS	
Animal I.D. #								· · ·		
34-1	11/2	11/8	14/3	159	24/13	0/0	11/12	0/0		
	R-2 E-2	R-2 E-1	R-3 E-3	R-3 E-3	R-2 E-2	R-00	R-3 E-3	E-0	-	
		u=40	4-60				4-60			
				·						
	J						0=1	not o	pporent	
All measurements in N/A = Not applica N/R = Not required	ble	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe	0	لىد AC سىر سىر مى ش	lcera ine s cont	tion of	gdose si reviously 38-96 pm	tes
- \ 1 \	,						Ψ <i>σ</i> .	10 25		
Site A <u>5-110</u>						//	ביה מסת	. سينراز	noted a	زره
Site B 25 - 2 60	سمحسا	- tate				بسراء			4 = small	_
Site C 5 2 107,	· HWi	-CHC	3						5= medin	m-1
Site D 25	٠	tata							6=lagu	
Site E 5_2 (07	lo Lim	CHCI:	1							
Site F 25.0 ba		atata		* . .						
Site G 1 - 2 ~	H to	7							•	
Site H 25-0-	<u> Lo</u>	سلصل								

Form No. MREF-LESION.S12-07
Appendix A

Reverse by (24) Th

Project #: G15	55- 38A	7				Date:	6-2	1-96		
MREF Protocol					-					
Day:	L	esion Re	ead By:		<u> </u>	esions R	Recorde	d By: 🔟	Omm	
Lesion Sites	А	С	E	G	В	D	F	Н	COMMENTS	
Animal I.D. #				,					4	
339	109	1813	12/12	9/10	0/6	19/14	00	28/3		
	E-2	R-3 E-3	F-3	R-3 E-2	R-0 E-0	R-2 E-2	E-0	R-2 E2		
		4-40	4=40	·						
										_
										_
	·]
										4
							2-1	10+00	20070	j
All measurements in		ers		Erythema	•	Λά	(- N	ر ما ما	parent doce frevious 62:-96	ر بين
N/A = Not applicat N/R = Not required			E = 1 =	Edema Mild	Ų	عد شاہرا مد	بلاوي	wet	Freviou	المالية المالية
				Moderate Severe		-	سمدين	diel	6 21-96	DAM
Site A 5-010	57× 77×	V: C+				u =	ulc	viati	مسكفاءر سن	رنعه ک
								4/-	- small	
Site B 25 L A							•	5=	wilden	i
Site C_5_10	lo Lin	-CHC	/3					6 =	large	
Site D_25_0 b	سے سے	tester								
Site E_5_010	OH of	C#4	تلء							
Site F_ 25_0 ~	4	atesta						٠	~•	
Site G_ l L ria	TH Z								•	
Site H_ 25,2 6	مسع	-55-	tram	-				Renzen	jed by	
Form No. MREF-LESIO	N.SIZ-07							C 7	الكامر د	124/36

Appendix A

Project #: G15	55- 38A					Date:	6-	21-96	_	
MREF Protocol			3	Stud	ly Direct	or: Car	l Olson			
									2	
Day:	L	esion Re	ead By:	$D_{\overline{A}}$	L	esions R	ecordeo.	d By:	CIRCA	_
Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS	
Animal I.D. #										
342	2214	11 8	9/10		23/15	00	11/22	00		
	R-3 E-3	R2 E-2	R-2 E-2	R=2 E=2	R-2 E-1	R-0	R-2 E-2	R-0		
]
·										
										}
				-						
	L		_				0-,	mot o	apparent	
All measurements in N/A = Not applica		ers		Erythema Edema		•			5 /	
N/R = Not require/			1 =	Mild Moderate						
			_	Severe						
Site A 5-210°	1. Li	~ C++	تاء				•			
Site B Salk							•			
										•
Site C 5-2107										
Site D 25.2 no	L mont	testre	-							
Site E_5_210	10 HW:	_CHC	EL							
Site F_25, 26	20	toetee		•						
Site G_lala	4 200	0						Rive	with by	
Site H_ 25.2	<u></u>	tester						<u>C</u> .	TOFA	6/24/96

Form No. MREF-LESION.SIZ-07

Project #: G15	55- 38A	7				Date:	<u>_6-3</u>	27-96	_	
MREF Protoco	1#: <u>10</u> 9	Pi.	<u> </u>	_ Stuc	ly Direc	tor: <u>Car</u>	l Olson			
Day:2	I	esion Re	ead By:	(A)	<u> </u>	esions F	Recorde	d By:	onini.	
Lesion Sites	А	С	E	G	В	D	F	Н	COMMENTS	
Animal I.D. #						*	1	4 6		_
340	12/1	8 8	11/14	11/14	19/3	19 R-1	22/6	00		_
	R-2 E-2	R-2 E-2	R-3	E-3	R-3 E-1	E-1	R-2 E1	R - 0		4
	4=41			4=60		-				4
										-
										-
										4
	ļ									.}
										1
								·		-
All measurements in N/A = Not applica N/R = Not required	ble	ers	E = 1 = 2 =	Erythema Edema Mild Moderate Severe		DACJ			ot appa	
Site A 540 107	6 HD	in C	HC-13			W	. سىد(metr	6-27-	TO BIMI
Site B <u> 25_11l .</u> .	<u>blue</u>	wad	estre	27						
Site C 524/0°	70 H.C	in C	HC-13		į	4-2-4			moted	ئنە ،
Site D 25 14 1	فطالما	Pacte	atria.	بر <u> </u>			-	Arria	Like you	
Site E <u>-5 Ml /C</u>	70 Li	-CHO	2/3				,	larg		
Site F 25 wk. L	ilui l	Jacte	ctra	ne.				•		**.
Site G / LLL M	int	HD								
Site H 25 ul	redu	acre.	Trea	ηW		Piu 2	سسيلا	لم و	7 Els-	L [25/56
Free No MOSS-1551	ח. כוס אח							`		

Form No. MREF-LESION.SIZ-07
Appendix A

Project #: G15	55- 38A	7				Date:	6, 0	27-96.	_	
MREF Protocol	1#: <u>10</u> 9	Ph	2cz 3	_ Stuc	ly Direct	tor: <u>Car</u>	-l Olsor			
Day:2	L	esion Re	ad By:	BH	L	esions F	Recorde	d By: <u>&</u>	IMAL	_
Lesion Sites	Α	С	E	G	В	D	F	Н	COMMENTS	
Animal I.D. #							.,		•	
345	9/3	19/14	11/10	14/15	00	19-15	21/5	2016		
3.75	R-3 C3	R-3	R-2 4-3	R-2 E-3	R-0 E-0	R-2	R-1	R-2 E-1		
	U-6	426	iL=b	ا سال				4=4		
	(1)	<u></u>	6	<i>(</i>)						
				•						
								·		
									•	
										,
									1-	4
All measurements in N/A = Not applica N/R = Not required	ble l		E = 1 = 2 = 3 =	Erythema Edema Mild Moderate Severe		DAC Sirl	ulci not p		ot oppor	
Site B <u>25 ul</u>) I —		. + '		
					_	~~,		ilrial	-notecle	w:
Site C <u>5ul/0</u>							•	midi		
Site D.25ull	ilun 1	Nost.	Thear	<i>,</i> ,,				large		
Site E <u>5ul 10</u>								•		
Site F <u> 25 ul</u> 1			Trea	m		•				
Site G/ul.m			•		Ω		1	(ala
Site H <u>25-11</u>	l-lue-1	wast	catrin	·m·	K	سالگدین	سرا ل	01	DPS- 6/2	2 [7.7-

Form No. MREF-LESION.SIZ-07

Project #: G155	55- 38 <i>A</i>	7				Date:	6.0	77-96	<u>-</u>	
MREF Protocol	#: 10	P	have 3	3_ Stud	dy Direct	or: <u>Car</u>	l Olson			
Day:	I	esion R	ead By:	_ B 13	L	esions R	Recorded	i By: _∠	CALAL	
Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS	
Animal I.D. #								,	_	
351	13/15	159	10/5		16/19	15/12	2/12	00		
	R-2 E-2	R-3	R-3 E-2	F-3 E-2	R-1 E-2	R-1 E-1	K-1	R-0 I0		
		4-4 ₀	11-Z	u-50						
										ı
			1							
All measurements in a N/A = Not applicable N/R = Not required	le		E = 1 = 2 = 3 =	Erythema Edema Mild Mioderate Severe		1AC	uéc vece		net appa net des portrois	
Site B 25 ul be			_	~		u =	ماس	er al	in note	نمما
Site C <u>5 ul 10 ° /</u>						,		= &N		
Site D <u>25 ul re</u>				.					diem	
Site E.5 ul 10%								la.	40	
Site F <u>25-12</u>				.YI~		-				
Site G/ul me	ati	HD								
Site H_25_ulx			strea	.wn		0	,	r 0	TORSL	6/25/9
Form No. MREF-LESION	.SIZ-07					Cerr	urch	النس ر	- 1 000	ί - ί -

Appendix A

Project #: G15	55- 38A	\				Date:	6.3	7-96	_
MREF Protocol	#: <u>10</u> 9	F-1	lace.	Stud	y Direc	tor: <u>Car</u>	l Olson		
Day:	L	esion Re	ad By:	RL	L	esions R	Recorded	By: 🔨	Jrepa
Lesion Sites	Α	С	E	G	В	D	F	Н	COMMENTS
Animal I.D. #						_			
3.52	119	12-9	11/12	13/15	00	16 15	15/12	15 9	
	R-2 E-2		R-3 C-3	E-3	R-0 E-0	R-1 E-2	R-1 E-1	£-2 €-i	
	il-50	2-4		u-60					
·									
					-				
								0=m	it apparent
All measurements in N/A = Not applicat		ers		Erythema Edema					-7-7
N/R = Not required			1 =						
				Severe	<i>(</i>)	54) /==	27-9	6 Rm	m
Site A Sul 107	6 HD	in CH	C/3						
200 11					2	AC U	lcero	tion s	of dose site

Site B 25 ul rid Wastatrian

Site C 5 ul 10% HN in CHC13

Site D 25 ul blue Wastestran

Site E 5 ul 10% Lin CHC13

Site F 25 ul red Wastestran

Site G / ul mest HD

Site H 25 ul blue Wastestran

Form No. MREF-LESION.SIZ-07

Appendix A

@ AC alceration of dose sites were not perviously recorded 6-27-98 DMM

4- ulceration noted as: 4- small 5- medium 6- large

Reviewed by CT Dem 6/25/86

	T	1	1	1	L		T	Γ	COMMENTS
Lesion Sites	A	С	E	G	В	D	F	H	COMMENTS
Animal I.D. #	10- 7	111 2	112/	10,	16	110/	116/	1	
383	150		10-18	10	P-1	R-15(2	1714 1K-i		
	₹-3 .43.23	5-3 5-3	g-2 E-2	-	<u>[= </u>	E-0	£-,		
			 						
· · ·			-						
					-				
				1				-	
	.1		1 =	Mild					
ite B/Oul le	% L		3 = {C13	Modera Severe	e				
ite A <u>1011</u> 10	% Li lu: ii	Jast	3 = 1C13 estres	Severe	e				
ite A <u>1011</u> 0	% L.: Vi Vo H C	Dast Dim C	3 = LC13 estres .HC13	Severe					
ite A <u>1011</u> 10 ite B <u>1011 b</u> ite C <u>1011 10</u> ite D <u>1011 Ch</u>	% Li lu: Vi % Ho arcos	Davi Duic Un b	3 = IC13 estrea IHC/3 esteo	Severe					
ite A 10 mg 10 ite B / O mg 10 ite C 10 mg 10 ite D 10 mg Ch ite E / O mg 10°	% Li % HC ar con	Dast Din C Din Ch	3 = IC13 Estrea HC13 esteci HC13	M treas				i?	
ite A <u>1011</u> 10 ite B <u>1011 b</u> ite C <u>1011 10</u> ite D <u>1011 Ch</u>	% Li % HC ar con	Dast Din C Din Ch	3 = IC13 Estrea HC13 esteci HC13	M treas				ىدەڭا	ward by C';

	Project #: GL	555- 38 <i>8</i>	<u>.</u>				Date:	8-1	4-96		
	MREF Protoco				_						
	Day:2_	L	esion Re	ead By:	Ab_	L	esions R	Recorded	1 By: _ <i></i>	<u>Cmm</u>	
	Lesion Sites	A	С	E	G	В	D	F	н	COMMENTS	
	Animal I.D. #					1.0	ســـــــــــــــــــــــــــــــــــــ	FII -	101/	7	
	385	13/5	12/8	12 1h	0	12-2 1R-2 E-2	13 R. 1 E-1	R-153	0		
		2-2 E-2	R-3 E-2	= 3		E-2	E-1	E-0			
				·							
		·									
		-									
										na not app	
	All measurements in N/A = Not applied N/R = Not require	able d		1 = 2 = 3 =	Edema Mild Moderate Severe						
	Site Aloud 10°										
	Site B 10 2 1										
	Site DIO ul				ฑส						
	Site E 10.110								Ø		9Q -
	Site F 10 C.							•	الف	iewally 97:	tlar.
an an	Site 0	<u> </u>									
	Appendix I Form No. MREF-LESI CAC Jhe	A 10N.SIZ-07	tes u	رىئىن	nat.	83 wed	on		tag s	3-13-96 Emr	~
	@IF 8-14-	76 Bn	im.	, ,	1.1656	ا ۴ د پ	مسلئل	· mine	e at-	ienela detue	سهرد

Day:	L	esion R	Read By:	23	I	Lesions I	Recorded	1 By: _	KMINC
Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS
Animal I.D. #									
400	17/7	15/5	1521	01		11-14		0	
•	という	R-3 53	R-3		R-150	R-1 E-0	K-1 E-1		
						ļ			
									<u> </u>
			1						·
				-					
All measurements in	millimete	ers	R =	•	a			o=m	et appare
N/A = Not applicab N/R = Not required	le		E = 1 = 2 = 3 =	Edema					ot appare
N/A = Not applicable N/R = Not required site A. Dull 10°	ile ZHI	Tinc	E = 1 = 2 = 3 = :HC/3	Edema Mild Moderat Severe				0=m	et appare
N/A = Not applicable N/R = Not required ite A. Dul 0° ite B 10 ul Cl	he To HI	Tinc I Wa	E = 1 = 2 = 3 = :HC/3	Edema Mild Moderat Severe				o=m	et appare
ite Blowl Ch	To HI	Jin C S Wa in Cl	E = 1 = 2 = 3 = :HCl3 :HCl3	Edema Mild Moderat Severe				o=m	et appare
ite B 10 ul 100 ite C 10 ul 100 ite D 10 ul 100 ul	To HI	Jin C S Wa in C!	E = 1 = 2 = 3 = :HCl3 :HCl3 HCl3	Edema Mild Moderat Severe				o=m	et appare
ite Double ite Double ite Double ite Double ite Double ite Double	To HI	Tinc O WO in Cl aster in C.H	E = 1 = 2 = 3 = HC/3 HC/3 Etran C/3	Edema Mild Moderat Severe				o=m	et appare
ite B 10 ul 100 ite C 10 ul 100 ite D 10 ul 100 ul	To HI	Tinc O WO in Cl aster in C.H	E = 1 = 2 = 3 = HC/3 HC/3 Etran C/3	Edema Mild Moderat Severe					
ite Double ite Double ite Double ite Double ite Double ite Double	To HI	Tinc O WO in Cl aster in C.H	E = 1 = 2 = 3 = HC/3 HC/3 Etran C/3	Edema Mild Moderat Severe					et appare

Lesion Sites	A	С	E	G	В	D	F	Н	COMMENTS
Animal I.D. #				7. —		111 /	111/	01	
389	13/3	12 31 R-3 E-3	121 	0	5 19 R5 Q	R I E I	F-0		
				1 /			<u> </u>		
ll measurements	in millime	eers		Erythe				C =	not appar
/A = Not applic /R = Not require te A / O u Q /	eable ed <u>07, HJ</u>	<i>کیسر</i>	E = 1 = 2 = 3 = 2HC/3	Edema Mild Modera Severe				C ==	not appar
A = Not applied $A = Not required$ the A $A = Not required$ the B $A = Not required$ the C $A = Not required$	ed 6	In Co astes in Ct	E = 1 = 2 = 3 = 2HC13 tream	Etlema Mild Modern Severe				C ==	net appar
A = Not applicate A Not require to B D L A te D D L A te D D D D D L A te D D D D D D D D D D D D D D D D D D	ed work Led w	sin Ct astes in Ct Daste	E= 1= 2= 3= CHC/3 Tream HC/3 Stream	Etlema Mild Modern Severe					not appar
A = Not applied $A = Not required$ the A $A = Not required$ the B $A = Not required$ the C $A = Not required$	ed word Led	linco astes in Ct Daste Din C	E= 1= 2= 3= 2HC/3 tream HC/3 Stream	Edema Mild Modern Severe	ate				

	<u> 1555- 384</u>	<u>A</u>				Date:	_8-	30-	96	<u></u>
MREF Protoc	ol#: 10	9 Pho	23	3	Study Direc	tor: <u>Ca</u>	<u>ri Oison</u>)		
Day:	I	esion R	ead By:		<u> </u>	Lesions I	Recorde	d By	:	Ric.
Lesion Sites	A	С	E	C	В	D	F	I	H	COMMENTS
Animal I.D. #							1	10.		
379	1149	13 10	11014	0	100	196	100	0		
	K-3	R-3	R-1		18-0	R-0	GO-00			
							1			
						1				
						1				
	-								_	
				- -	-	!	1			
All measurements	in millimete		R =	Frythe	-ma		0=	L net	<u></u>	wrent
All measurements in N/A = Not application N/R = Not requires	ed		E = 1 1 = 1 2 = 1 3 = 3	Erythe Edema Mild Moder Severe	ate .		O=	L net	29	porent
N/A = Not applic N/R = Not require Site A_Jul 107	eable ed	CHC13	E = 1 1 = 1 2 = 1 3 = 5	Edemi Mild Moder Severe	ate		0=	Not.	29	porent
N/A = Not applic N/R = Not require Site A Jul 107 Site B 25xl C	table ed To Lin	CHC13	E = 1 = 1	Edemi Mild Moder Severe	ate			,	·	porent
N/A = Not applie N/R = Not require Site A Jul 107 Site B 25xl C Site C Jul 106	Larco	CHC13 of War	E = 1 1 = 1 2 = 1 3 = 5	Edema Mild Moder Severe	ate	Res		: IL	`	•
N/A = Not applic N/R = Not require Site A Jul 107 Site B 25xl C	Larco	CHC13 of War	E = 1 1 = 1 2 = 1 3 = 5	Edema Mild Moder Severe	ate	Revi	0= ()	: IL	`	•
N/A = Not applie N/R = Not require Site A 5 ul 107 Site B 25 ul 107 Site C 5 ul 107	Larco	CHC13 cl War al War	E = 1 = 1 2 = 1 3 = 5 itestre Cl3	Edema Mild Moder Severe	ate	Resi		: IL	`	prest 91301
N/A = Not applic N/R = Not require Site A Jul 107 Site B 25x1 C Site C Jul 104 Site D 25x1 C	Jacon Larcon Larcon Larcon Larcon Larcon	CHC13 ch War al war	E=1 1=1 2=1 3=3 testre	Edermi Mild Moder Severe	ate	Revi		: IL	`	•
N/A = Not applie N/R = Not require Site A 5 ul 107 Site B 25 ul 107 Site C 5 ul 107 Site D 25 ul 107 Site E 5 ul 107	Jacon Larcon Larcon Larcon Larcon Larcon	CHC13 ch War al war	E=1 1=1 2=1 3=3 testre	Edermi Mild Moder Severe	ate	Revi		: IL	`	•
N/A = Not applie N/R = Not require Site A Jul 107 Site B 25 L C Site C Jul 106 Site E Jul 107 Site E Jul 107 Site F 25 L C	Jacon Larcon Larcon Larcon Larcon Larcon	CHC13 ch War al war	E=1 1=1 2=1 3=3 testre	Edermi Mild Moder Severe	ate	Revi		: IL	`	•

MREF Protocol #: 109 Place 3.3 Study Director: Carl Olson Day: Z Lesion Read By: CTO Lesions Recorded By: Protocol #: Lesion Sites A C E G B D F H COMMENTS Animal I.D. # 380 10 10 3 0 9 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	Project #: <u>G1555-38</u>	<u>A</u> .				Date:	8-3	30-90	<u>-</u>	
Lesion Sites A C E G B D F H COMMENTS Animal I.D. # 380 196 170 3 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0 0	MREF Protocol #: 11	19 P2	102.3	3_ Stu	dy Direc	tor: <u>Ca</u>	d Olson			
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APPENDIX B

Dosage Site Code and Histopathology

Definitions Used in Histopathologic Evaluations and an Explanation of the Grading of Lesion Severity

Microblister: Loss of epidermal basal cell attachment to the underlying basement membrane of at least two adjacent cells. The loss of attachment creates a space which may appear empty, full of proteinaceous fluid, or filled with neutrophils. One or a few isolated small areas of detachment is graded 1, minimal. Many such areas of detachment, or several larger (10 or more contiguous cells) areas of detachment is graded 2, mild. When half or more of the epidermis in the tissue section is detached from the dermis, it is graded 3, moderate. Such lesions typically have a much larger space between the basal cells and the dermis. When nearly all of the epidermis is separated from the dermis, it is graded 4, marked. In such situations, there are usually focal, point attachments, so the entire epidermis is not lifted along the full width of the section.

Epidermal necrosis: The epidermal cells exhibit cytoplasmic eosinophilia, nuclear loss or pyknosis, and are generally shrunken. If only individual cells are affected, it is graded 1 (these are generally isolated basal cells). If small areas are affected, with normal areas in close proximity, it is graded 2. If the epidermis exhibits cell death in a full-thickness (all layers of epidermis) pattern, and affects half or more of the skin section, it is graded 3. If the epidermis is virtually entirely necrotic, it is graded 4. Severe ulcers assume that the epidermis is necrotic.

Follicular necrosis: If isolated epithelial cells of the hair follicles exhibit eosinophilia or pyknosis, it is graded 1. If clusters of adjacent cells within follicles are dead, it is graded 2. If cells of half or more of a particular hair follicle are dead, it is graded 3. Grade 4 lesions have complete necrosis of the follicular epithelium underlying much of the epidermal lesion area. This indicates that the agent has penetrated deeply.

Dermal necrosis: Loss of collagen fiber integrity, evidenced by pale eosinophilic staining and homogeneous appearance, indicates necrosis of dermal fibers. With only isolated areas, it is graded 1. Multiple areas are graded 2. Necrosis of most of the superficial dermal collagen in the lesion area is graded 3. A grade four lesion requires deep (to the base of the associated adnexa) dermal necrosis.

Hemorrhage: Extravasated erythrocytes is hemorrhage. A few isolated foci is graded 1. Multiple, common foci is graded 2. Large pools of blood is graded 3. A grade four lesion requires a massive area of blood pooling with displacement of large areas of dermal collagen.

Vascular necrosis: Loss of integrity of a medium to large blood vessel is vascular necrosis. Grading depends upon the number of vessels affected and the severity. Partial necrosis of one vessel is graded 1 to 2. Complete necrosis of a vessel is graded 3; multiple such lesions are graded 4.

Pustular epidermitis: Collections of neutrophils in the epidermis proper is graded by extent; one or two small foci is graded 1; three or more small foci is graded 2; one or more large foci is graded 3; a grade four lesion would indicate massive infiltration of the entire epidermis by neutrophils.

Task 95-38, Phase 2a, Day 1

Key for HGPs #301 and 305 dosed 2/19/1996. Exposure duration - 2 hr.

Animal # 301

Site	Treatment
A	10 μL of 10% HD in CHCl ₃
В	50 μL of 10% HD in CHCl ₃
С	10 μL of 10% HN in CHCl ₃
D	50 μL of 10% HN in CHCl ₃
E	10 μL of 10% L in CHCl ₃
F	50 μL of 10% L in CHCl ₃
G	1 μL of neat HD
H	·

Site	Treatment
A	10 μL of 10% HN in CHCl ₃
В	50 μL of 10% HN in CHCl ₃
С	10 μL of 10% L in CHCl ₃
D	50 μL of 10% L in CHCl ₃
E	10 μL of 10% HD in CHCl ₃
F	50 μL of 10% HD in CHCl ₃
G	1 μL of neat HD
Н	

Dosing Date: 2/19/96

MREF Task 95-38 G1555-38A

Animal # 301	Site	A	В	С	D	E	F	G		
Histopathology Marker	Histopathology Markers:									
Microblister		2	2	2	2	3	4	2		
Epidermal Necrosis		2	4	4	3	3	4	3		
Follicular Necrosis		3	4	4	4	2	4	4		
Dermal Necrosis		0	0	0	0	0	0	0		
Vascular Necrosis		0	0	0	0	0	0	0		
Hemorrhage		0	0	0	0	1	2	0		
Pustular Epidermitis		0,	0	0	0	0	0	0		
Notes: all lesions are centrally located; some normal skin presentall	nt on		mild dermal inflam	min dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam		

Animal # 305	Site	A	В	С	D	E	F	G
Histopathology Marker	s:							
Microblister		2	2	3	3	3	2	2
Epidermal Necrosis		4	4	4	4	4	4	4
Follicular Necrosis		3	4	4	4	4	4	4
Dermal Necrosis		0	0	0	0 .	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	1	0	1	1	0
Pustular Epidermitis		0	0	0	0	0	0	0
Notes: all lesions are centrally located; some normal skin presentall	nt on	mild dermal inflam	mild dermal inflam	mild dermal inflam		mild dermal inflam	min dermal inflam	mild dermal inflam

Degree of Severity Grading Scale:

14

^{0 =} Normal, 1 = Minimal, 2 = Intermediate, 3 = Moderate, 4 = Severe
Appendix B 94

Task 95-38, Phase 2a, Day 2

Key for HGPs #306 and 309 dosed 2/21/1996. Exposure duration - 2 hr.

Animal # 306

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	10 μL of 10% L in CHCl ₃
С	5 μL of 10% HD in CHCl ₃
D	10 μL of 10% HD in CHCl ₃
E	5 μL of 10% HN in CHCl ₃
F	10 μL of 10% HN in CHCl ₃
G	1 μL of neat HD
Н	

Site	Treatment
A	10 μL of 10% HD in CHCl ₃
В	5 μL of 10% HD in CHCl ₃
С	10 μL of 10% HN in CHCl ₃
D	5 μL of 10% HN in CHCl ₃
E	10 μL of 10% L in CHCl ₃
F	5 μL of 10% L in CHCl ₃
G	1 μL of neat HD
Н	

Animal # 306	Site	A	В	С	D	E	F	G	
Histopathology Markers:									
Microblister		4	3	1***	3	4	2***	1***	
Epidermal Necrosis		4	4*	4***	4	4	4***	4***	
Follicular Necrosis		4	4	4	4	4	4	4	
Dermal Necrosis		1	1**	2	0	0	2	2	
Vascular Necrosis		0	0	0	0	0	0	0	
Нетогтнаде	٠.	1	0	0	0	0	0	0	
Pustular Epidermitis		0	0	0	0	0	0	0	
Notes: *focal ulceration **deep dermal edema ***large ulcer preclude much blister potential	s	mod dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	

Animal # 309 Site		A	В	С	D	Е	F	G
Histopathology Marker	s:							
Microblister		3	0*	4	4	4	4	3
Epidermal Necrosis		3	4*	4	4	4	4	4
Follicular Necrosis		4	4	3	2	4	3	4
Dermal Necrosis		1	2	0	0	0**	0	0
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	1	0	0
Pustular Epidermitis		1	0	1	1	0	0	0
Notes: *large ulceration preclu blister potential **deep dermal edema	d e s	mild dermal inflam	mild dermal inflam	mod dermal inflam	mod derm infla m	mod dermal inflam	mod dermal inflam	mild derm infla m

Note: Some normal skin is present on all sections, both animals; lesions are centrally located in trimmed area.

Degree of Severity Grading Scale:

Task 95-38, Phase 2a, Day 3

Key for HGPs #312 and 316 dosed 2/27/1996. Exposure duration - 1 hr.

`Animal # 312

Site	Treatment
A	10 μL of 10% L in CHCl ₃
В	5 μL of 10% L in CHCl ₃
·c	10 μL of 10% HD in CHCl ₃
D	5 μL of 10% HD in CHCl ₃
E	10 μL of 10% HN in CHCl ₃
F	5 μL of 10% HN in CHCl ₃
G	1 μL of neat HD
H	

Site:	Treatment
A	10 μL of 10% HN in CHCl ₃
В	5 μL of 10% HN in CHCl ₃
С	10 μL of 10% L in CHCl ₃
D	5 μL of 10% L in CHCl ₃
E	10 μL of 10% HD in CHCl ₃
F	5 μL of 10% HD in CHCl ₃
G	1 μL of neat HD
Н	

Animal # 312	Animal # 312 Site		В	С	D	E	F	G	
Histopathology Markers:									
Microblister		3	3	3	3	4	3	3	
Epidermal Necrosis		4	4	4	4	4	4	4	
Follicular Necrosis		4	4	4	4	4	3	4	
Dermal Necrosis		0*	0*	0	0**	0	0	0*	
Vascular Necrosis		0	0	0	0	0	0	0	
Hemorrhage		1	2	0	0	0	0	0	
Pustular Epidermitis		0	0	0	0	1	2	0	
Notes: *mod dermal edema **minimal dermal edema		mild dermal inflam	mod dermal inflam	mild dermal inflam	mild derm infla m	mild dermal inflam	mod derm inflam	mild derm infla m	
Animal # 316	Site	A	В	С	D	Е	F	G	
Histopathology Marker	s:							-	
Microblister		3	4	4	4	3	3 .	3	
Epidermal Necrosis		4	4	4 .	4	4	4	4	
Follicular Necrosis		4	3	4	4	4	4	4	
Dermal Necrosis		0*	0	0**	0**	0	1	0**	
Vascular Necrosis		0	0	0	0	0	0	0	
Hemorrhage		0	0	2	2	1	0	0	
Pustular Epidermitis		1	1	0	0	1	1	2	
Notes: *minimal dermal edema **moderate dermal edema		mod dermal inflam	mod dermal inflam	mod dermal inflam	sever e derm infla m	mild dermal inflam	mod dermal inflam	mod derm infla m	

Note: All sections (312 and 316) have normal, unaffected skin at one or both margins of the section.

Degree of Severity Grading Scale:

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

Allen W. Singer, D.V.M. Appendix B

Task 95-38, Phase 2b, Day 1

Key for HGPs #311, 313, 315, 317, and 324 dosed 3/5/1996. Exposure duration - 1 hr.

Animal #311

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% L in CHCl ₃
D	20 μL of neutralization solution
Е	5 μL of 10% HD in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
Н	20 μL of neutralization solution

Site	Treatment			
A	5 μL of 10% HD in CHCl ₃			
В	20 μL of neutralization solution			
С	5 μL of 10% HN in CHCl ₃			
D	20 μL of neutralization solution			
Е	5 μL of 10% L in CHCl ₃			
F	20 μL of neutralization solution			
G	1 μL of neat HD			
Н	20 μL of neutralization solution			

Animal #315

Site	Treatment
·A	5 μL of 10% HN in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% L in CHCl ₃
D	20 μL of neutralization solution
E	5 μL of 10% HD in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
H	20 μL of neutralization solution

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% HD in CHCl ₃
D	20 μL of neutralization solution
Е	5 μL of 10% HN in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
н	20 μL of neutralization solution

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	20 μL of neutralization solution
С	5 μL of 10% HN in CHCl ₃
D	20 μL of neutralization solution
E	5 μL of 10% L in CHCl ₃
F	20 μL of neutralization solution
G	1 μL of neat HD
Н	20 μL of neutralization solution

Animal # 311 Site		Α	В	С	D	Е	F	G	Н
Histopathology Marke	Histopathology Markers:								
Microblister		3	0	3	0	2	0	3	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		2	0	4	0	4	0	4	0
Dermal Necrosis		0	0	0	0	0	0	0*	0
Vascular Necrosis		0	0	. 0	0	0	0	0	0
Hemorrhage		2	0	3	0	0	0	1	0
Pustular Epidermitis		2	0	0	0	1	0	0	0
Note: *moderate deep dermal edema		mod dermal inflam	·	mod dermal inflam		mod dermal inflam		mod derm infla m	

Animal # 313	Site	A	В	С	D	E	F	G	Н
Histopathology Mark	Histopathology Markers:								
Microblister		3	0	4	. 0	4	0	2	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		4	0	4	0	3	0	4	0
Dermal Necrosis		0	0	1	0	0	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Нетоппаде		0	0	0	0	1	0	0	0
Pustular Epidermitis		0	0	0	0	0	0	0	0
Note: *moderate deep dermal edema)	mild derm inflam		mod dermal inflam	min dermal inflam	mod dermal inflam		mild dermal inflam	

Animal # 315	Site	Α	В	С	D	Е	F	G	Н
Histopathology Market	Militar # 22								
Microblister		2	0	4	0	3	0	2	0
Epidermal Necrosis		3	0	4	0	4	0	4	0
Follicular Necrosis		2	0	4	0	4	0	4	0
Dermal Necrosis		0	0	1	0	1	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		0	0	1	0	2	0	0	0
Pustular Epidermitis		1	0	0	0	0	0	0	0
Note: *moderal derma	1	mod dermal inflam		marke d dermal inflam		mod dermal inflam		mild dermal inflam	

Animal # 317 Site		A	В	С	D	Е	F	G	H
Histopathology Mar	kers:								
Microblister		2	0	2	. 0	3	0	2	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		4	0	4	0	3	0	4	0
Dermal Necrosis		0*	0	2**	0	0	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		2	0	1	0	0	0	0	0
Pustular Epidermitis	;	0	0	0	0	2	0	0	0
Notes: *mild dermal edema **focal ulceration(s)		mild dermal inflam	min dermal inflam	mod dermal inflam		mod dermal inflam		mild dermal inflam	

E-13

Animal # 324	Site	Α	В	С	D	E	F	G	Н
Hîstopathology Marke	Histopathology Markers:								
Microblister		4	0	4	0	4	0	3	0
Epidermal Necrosis		4	0	4	0	4	0	4	0
Follicular Necrosis		4	0	2	0	4	0	4	0
Dermal Necrosis		1	0	0	0	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Нетоггнаде		0	0	0	0	1	0	0	0
Pustular Epidermitis		0	0	1	0	0	0	0	0
Notes:		mod dermal inflam	min dermal inflam	mod dermal inflam		mod dermal inflam		min dermal inflam	

Note: Normal (unaffected) skin present laterally on all sections where lesions were observed.

Histopathological Markers
Degree of Severity Grading Scale
DVM

3/7/96 Allen W. Singer,

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

Task 95-38, Phase 3, Day 1

Key for HGPs #310, 491, 493, and 498 dosed 3/13/1996. Exposure duration - 1 hr.

Animal #310

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Blue waste stream
Е	5 μL of 10% L in CHCl ₃
F	25 μL of Charcoal waste stream
G	1 μL of neat HD
H	

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Red waste stream
Е	5 μL of 10% HN in CHCl ₃
F	25 μL of Blue waste stream
· G	1 μL of neat HD
Н	

E-15

Site	Treatment
Α	5 μL of 10% HD in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HN in CHCl ₃
. D	25 μL of Blue waste stream
Е	5 μL of 10% L in CHCl ₃
F	25 μL of Charcoal waste stream
G	1 μL of neat HD
H	

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Charcoal waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
H	

Animal # 310	Site	Α	В	С	D	E	F	G
Histopathology Markers:								
Microblister		2	0	4	2	1	0	1
Epidermal Necrosis		4	1	4	4	4*	2	4*
Follicular Necrosis		4	0	4	1	4	0	4
Dermal Necrosis		0	0	1	0	3	0	3**
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	0	0	1
Pustular Epidermitis		0	1	0	1	0	1	0
Notes: *marked ulceration **moderate dermal edema		mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mild dermal inflam	mod dermal inflam
Animal # 491	Site	A	В	С	D	E	F	G
Histopathology Mar	kers:							
Microblister		4	0	1	0	4	3	2
Epidermal Necrosis		4*	1	4**	0	4	4	4
Follicular Necrosis		4	0	4	0	3	0	4
Dermal Necrosis		3	0	3	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage	2	0	0	0	0	0	0	
Pustular Epidermitis		0	0	0	0	0	0	0
Notes: *mild ulcerat **marked ulceration	ion	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam

Animal # 493	Site	A	В	C	D	E	F	G	
Histopathology Markers:									
Microblister		1*	0	4	4	2	. 0	2*	
Epidermal Necrosis		4**	0	4	4	4**	1	4**	
Follicular Necrosis		4	0	3	0	4.	0	4	
Dermal Necrosis		3	0	0	0	3	0	3	
Vascular Necrosis		0	0	0	0	0	0	0	
Hemorrhage		0	0	0	0	1	0	0	
Pustular Epidermitis		0	1	0	0	0	0	0	
Notes: *at edge of uld **marked ulceration	cer	mod dermal inflam							

Animal # 498	Site	Α	В	С	D	E	F	G		
Histopathology Markers:										
Microblister		2*	3	. 3	0	3	0	1		
Epidermal Necrosis		4**	4***	4***	0	4**	0	4***		
Follicular Necrosis	Follicular Necrosis		0	4	Ö	4	0	4		
Dermal Necrosis		3	1 .	2	0.	3	0	2		
Vascular Necrosis		0	0	0	0	0	0	0		
Hemorrhage	Hemorrhage		0	1	0	0	0	0		
Pustular Epidermitis			0	0	1	1	0	0		
Notes: *at edge of u **marked ulceration ***minimal ulceration	1	mod dermal inflam	mild dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mild dermal inflam	mild dermal inflam		

Histopathological Markers: Degree of Severity Grading Scale 3/18/96
0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe Allen W. Singer, DVM

 $\mathcal{J}_{\mathcal{F}}$

Task 95-38, Phase 3, Day 2

Key for HGPs #494, 496, 497, and 499 dosed 3/21/1996. Exposure duration - 1 hr.

Animal # 494

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Red waste stream
E	5 μL of 10% HN in CHCl ₃
F	25 μL of Blue waste stream
G	1 μL of neat HD
Н	

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Blue waste stream
Е	5 μL of 10% L in CHCl ₃
F	25 μL of Charcoal waste stream
G	1 μL of neat HD
Н	

E-19

Site	Treatment
Α	5 μL of 10% HN in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Charcoal waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
H	

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Charcoal waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
Н	

Animal # 494	Site	A	В	С	D	Е	F	G	
Histopathology Markers:									
Microblister		4	0	1	0	3	2	3	
Epidermal Necrosis		4	0	4**	0	4	2	4***	
Follicular Necrosis		4	0	4	0	4	0	4	
Dermal Necrosis		0*	0	3	0	0	0	2	
Vascular Necrosis		0	0 ,	0	0	0	0	0	
Hemorrhage		3	0	0	0	0	0	0	
Pustular Epidermitis		0	0	1	0	1	0	0	

Animal # 496	Site	A	В	С	D	E	F	G		
Histopathology Markers:										
Microblister		0	0	0	4	1	0	2		
Epidermal Necrosis		4*	0	4*	3	4*	1	4*		
Follicular Necrosis		4	0	4	0	4	0	4		
Dermal Necrosis	Dermal Necrosis		0	3	0	4	0	3**		
Vascular Necrosis		0	0	0	0	0	0	0		
Hemorrhage	Hemorrhage		0	0	0	0	0	0		
Pustular Epidermitis	3	0	0	0	0	0	0	0		
Notes: *marked ulc precludes potential blister **mild dermal eden		mod dermal inflam	min dermal inflam	mod dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam		

Animal # 497	Site	A	В	С	D	E	F	G
Histopathology Mark	cers:							
Microblister		1	2	4	0	2	0	2
Epidermal Necrosis		4*	4	4	1***	4*	0	4*
Follicular Necrosis		4	0	4	0	4	0	4
Dermal Necrosis		3	0	0**	0	2	0	2**
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	0	0	0
Pustular Epidermitis		0	0	0	0	0	0	0
moderate dermal e	Notes: *marked ulceration **moderate dermal edema *mild epithelial cell edema			mod al derm n inflar	al derm	al dermal	mild dermal inflam	mod dermal inflam
Animal # 499	Site	A	В	С	D	E	F	G
Histopathology Mark	ers:							
Microblister		4	2	3	0	4	0	3
Epidermal Necrosis		4	3	4	2	4	0	4
Follicular Necrosis		4	0	4	0	4	0	4
Dermal Necrosis		0	0	2*	0	2	0	1*
Vascular Necrosis		0	0	0	0	0	0	0
Нетогтнаде	Hemorrhage		0	1	. 0	0	0	0
Pustular Epidermitis		1	0	0	0	1	0	0 ·
Note: *mild dermal ed	iema	mod dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mild dermal inflam

Histopathological Markers: Degree of Severity Grading Scale 3/25/96 0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe Allen W. Singer, DVM

Task 95-38, Phase 3, Day 3

"Fresh" Blue and Red waste streams received 6/19/1996

Key for HGPs #339, 341, 342, and 346 dosed 6/20/1996. Exposure duration - 1 hr.

Animal # 339

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Blue waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
H	25 μL of Blue waste stream

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Red waste stream
Е	5 μL of 10% L in CHCl ₃
· F	25 μL of Blue waste stream
G	1 μL of neat HD
Н	25 μL of Red waste stream

Site	Treatment
Α	5 μL of 10% L in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Red waste stream
E	5 μL of 10% HN in CHCl ₃
F	25 μL of Blue waste stream
G	1 μL of neat HD
н	25 μL of Red waste stream

Site	Treatment
Α	5 μL of 10% L in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HD in CHCl ₃
.D	25 μL of Blue waste stream
Е	5 μL of 10% HN in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
Н	25 μL of Blue waste stream

Animal # 339	Site	Α	В	С	D	E	F	G	Н
Histopathology Markers:									
Microblister		3	0	4	3	3	0	2	2
Epidermal Necrosis		4	0	4	4	4**	0	4	2
Follicular Necrosis		4	0	4	0	4	0	4	0
Dermal Necrosis		0	0	2*	. 0	2	0	0*	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	0	0	0	0
Pustular Epidermitis		1	0	1	1	1	0	0	0
Notes: *moderate dermal edema **focal ulceration(s)		mod dermal inflam	min dermal inflam	mod dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam	mild dermal inflam	min dermal inflam
									·
Animal # 341	Site	A	В	С	D	Е	F	G	H
Histopathology Marke	rs:								
Microblister	2	2	2	. 0	3	0	2	0	
		-			1				

Animal # 341 Site		A	В	. C	D	E	F	G	H
Histopathology Marker	Histopathology Markers:								
Microblister		2	2	2	. 0	3	0	2	0
Epidermal Necrosis		4*	4	4*	0	4*	4*	4*	0
Follicular Necrosis		4	1	4	0	4	2	2	0
Dermal Necrosis		3	1	2	0	3**	3	3**	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		1	0	0	0	1	0	1	0
Pustular Epidermitis		0	0	0	0	0	0	0	0
Notes: *focal ulceration(s); **moderate dermal edema		mild dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	min dermal inflam

Animal # 342	Site	Α	В	С	D	E	F	G
Histopathology Markers:								
Microblister		3	1	3	0	4	3	3
Epidermal Necrosis		4	4	4	0	4	4	4
Follicular Necrosis		4	0	4	0	4	1	4
Dermal Necrosis		0*	0	0*	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0	0
Hemorrhage		0	0	0	0	0	0	0
Pustular Epidermitis		0	0	1	0	0	0	0
Notes: *mild to moderate dermal edema	è	mild dermal inflam	min dermal inflam	mild dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mild dermal inflam

Animal # 346	Site	A	В	С	D	E	F	G	
Histopathology Markers:									
Microblister		2	0	2	1	4	0	2	
Epidermal Necrosis		4	0	4	4	4	0	4	
Follicular Necrosis		4	0	4	1	4	0	4	
Dermal Necrosis		0*	0	0	0	2	0	0*	
Vascular Necrosis		0	0	0	0	0	0	0.	
Hemorrhage		0	0	0	0	0	0	0	
Pustular Epidermitis		0	0	0	0	0	0	0	
Notes: *moderate dermal edema; **most of surface epithelium artifactually stripped away		mild dermal inflam	min dermal inflam	mod dermal inflam	mild dermal inflam	mild dermal inflam		mild dermal inflam	

Note: Normal (unaffected) skin presented laterally on all skin sections with lesions.

Histopathological Markers

6/25/96

Degree of Severity Grading Scale

Allen W. Singer, DV

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

Appendix B

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Task 95-38, Phase 3, Day 4

[&]quot;Fresh" Blue and Red waste streams received 6/19/1996

E-26

Key for HGPs #340, 345, 351, and 352 dosed 6/26/1996. Exposure duration - 1 hr.

Animal # 340

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Red waste stream
E	5 μL of 10% L in CHCl ₃
F	25 μL of Blue waste stream
G	1 μL of neat HD
H	25 μL of Red waste stream

Site	Treatement
Α	5 μL of 10% L in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Blue waste stream
E	5 μL of 10% HN in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
H	25 μL of Blue waste stream

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μL of Blue waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Red waste stream
E	5 μL of 10% HD in CHCl ₃
F	25 μL of Blue waste stream
G	1 μL of neat HD
н	25 μL of Red waste stream

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	25 μL of Red waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Blue waste stream
Е	5 μL of 10% L in CHCl ₃
F	25 μL of Red waste stream
G	1 μL of neat HD
Н	25 μL of Blue waste stream

E-28

Animal # 340	Site	Α	В	С	D	E	F	G	H
					<u> </u>				
Histopathology Ma	IKCIS.			г	1				_
Microblister		2*	3	3	0	3	3	0	0
Epidermalal Necro	sis	4**	4	4	0	4	4	4	1
Follicular Necrosis		4	2	4	0	4	0	4	0
Dermal Necrosis		2	0	1	0	0***	0	3***	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		2	0	0	0	2	0	2	0
Pustular Epidermi	tis	0	0	0	0	0	0	0	0
Notes: *at edge of **mild ulceration		mod dermal inflam	mild dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	min dermal inflam	min dermal inflam

Animal # 345	Site	Α	В	С	D	E	F	G	H
Histopathology Mar	Histopathology Markers:								
Microblister		3*	0	.2	1	1	0	1	2
Epidermal Necrosis	3	4**	0	4	4	4**	0	4	4
Follicular Necrosis		3	0	4	1 .	4	0	4	1
Dermal Necrosis		3	0	0	0	3	0	2***	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		2	0	1	0	2	0	1	0
Pustular Epidermit	is	0	1	0	0	0	0	0	0
Notes: *at one edge ulcer **marked ulceration present	of	mod dermal inflam	min dermal inflam	mild dermal inflam	mild dermal inflam	mod dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam

Animal # 351	Site	Α	В	С	D	Е	F	G	Н
Histopathology Mark	kers:						•		
Microblister		1	2	4	0	1*	2	1	0
Epidermal Necrosi	s	4	4	4	0	4**	4	4	0
Follicular Necrosis		4	1	3	0	4	1	4	0
Dermal Necrosis		0	0	0	0	3	0	3	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		0	0	1	0	1	0	1	0
Pustular Epidermiti	.s	0	0	0	0	0	0	0	0
Notes: *at one edge oulcer **marked ulceration present	of	mild dermal inflam	min dermal inflam	mod dermal inflam		mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam

Animal # 352	Site	Α	В	С	D	E	F	G	H
Histopathology Ma	arkers:								
Microblister		1*	0	2	1	3	0	2	2
Epidermal Necro	sis	4**	0	4**	4	4	0	4**	4
Follicular Necrosis	3	4	0	3	0	4	0	4	1
Dermal Necrosis		2	0	1	0	0***	0	3	0
Vascular Necrosis		0	0	0	0	0	0	0	0
Hemorrhage		2	0	1	0	1	0	1	0
Pustular Epidermi	tis	0	0	1	1	0	0	0	0
Notes: *at edge of **moderate ulcerat ***mild dermal e	ion	mod dermal inflam	min dermal inflam	mod dermal inflam	mod dermal inflam	mild derma l inflam		mod dermal inflam	mild dermal inflam

Histopathological Markers: Degree of Severity Grading Scale 7/1/96

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe Allen W. Singer, DVM

Appendix B 120

Task 95-38, Phase 3, Day 5

Blue and Red waste streams received 11/28/1995; Charcoal waste stream received 1/25/96.

E-30

Equal volumes of waste streams and 10% HD, HN and L solutions - 10 μ L

Key for HGPs #383, 385, 389, and 400 dosed 8/13/1996. Exposure duration - 1 hr.

Animal # 383

Site	Treatment	
A	10 μL of 10% L in CHCl ₃	
В	10 μL of Blue waste stream	
С	10 μL of 10% HD in CHCl ₃	
D	10 μL of Charcoal waste stream	
E	10 μL of 10% HN in CHCl ₃	
F	10 μL of Red waste stream	

Site	Treatment
A	10 μL of 10% HN in CHCl ₃
В	10 μL of Red waste stream
С	10 μL of 10% L in CHCl ₃
D	10 μL of Blue waste stream
E	10 μL of 10% HD in CHCl ₃
F	10 μL of Charcoal waste stream

Site	Treatment
A	10 μL of 10% HN in CHCl ₃
В	10 μL of Red waste stream
С	10 μL of 10% L in CHCl ₃
D	10 μL of Blue waste stream
E	10 μL of 10% HD in CHCl ₃
F	10 μL of Charcoal waste stream

Animal # 400

Site	Treatment
A	10 μL of 10% HD in CHCl ₃
В	10 μL of Charcoal waste stream
С	10 μL of 10% HN in CHCl ₃
D	10 μL of Red waste stream
E	10 μL of 10% L in CHCl ₃
F	10 μL of Blue waste stream

-5.5

E-32

MREF Task 95-38 G1555-38A

Animal # 383	Site	A	В	С	D	E	F
Histopathology Markers:							,,
Microblister		3	0	2	0	3	0
Epidermal Necrosis		4	0	4	0	4	0
Follicular Necrosis		4	0	4	0	3	0
Dermal Necrosis		0*	0	0	0	0	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage		2	0	1	0	0	0
Pustular Epidermitis		0	1	0	0	1	0
Notes: *moderate dermal edema		mod dermal inflam	·	mild dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam
Animal # 385	Site	A	В	С	D	E	F
Histopathology Markers:							
Microblister		4	0	4	1	3	0
Epidermal . Necrosis		4	0	4	1**	4	0
Follicular Necrosis		4	0	4	0	4	0
Dermal Necrosis		1	0	0*	0	0*	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage		0	0	2	0	0	0
Pustular Epidermitis		1	0	0	0	0	0
Notes: *mod dermal ed	ema	marked	min	mod		mild	min

Note: Some normal (unaffected) skin present at one or both ends of each section where lesions were present.

dermal

inflam

dermal

inflam

Histopathological Markers: Degree of Severity Grading Scale 0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

dermal

inflam

8/19/96 Allen W. Singer, DVM

dermal

inflam

dermal

inflam

**vacuolar degeneration of

epith cells leading to intra- and subepithelial microblister

ш-33 MREF Task 95-38 61555-38a

Animal # 389 Site		A	В	С	D	E	F
Histopathology Markers							
Microblister	3	0	2	2	2	0	
Epidermal Necrosis		4	0	4	2	4	1
Follicular Necrosis		2	0	4	1	4	0
Dermalal Necrosis		0	0	0*	0	0*	0
Vascular Necrosis		0	0	0	0	0	0
Hemorrhage		1	0	3	0	2	0
Pustular Epidermitis		1	0	0	0	0	1
Notes: *severe dermalal e	edema	mod dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam	mild dermal inflam
Animal # 400	Site	A	В	C	D	E	F
Histopathology Markers:							
Microblister		3	0	4	0	3	3
Epidermal Necrosis		4	0	4	0	4	2
Follicular Necrosis		4	0	2	0	4	1
Dermal Necrosis		0*	0	0*	0	0**	. 0
Vascular Necrosis		0	0	0	0	1	0
Hemorrhage		0	0	1	0	3	1
Pustular Epidermitis		1	0	1	·o	0	0
Notes: *mild dermal edema **severe dermal edema	ma	mod dermal		mod dermal	min dermal	mod dermai	mild dermal

Note: Some normal (unaffected) skin present at one or both ends of each section where lesions were present.

inflam

inflam

inflam

Histopathological Markers: Degree of Severity Grading Scale 0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

inflam

8/19/96 Allen W. Singer, DVM

inflam

Task 95-38, Phase 3, Day 6

"Fresh" Charcoal waste stream received 8/29/96.

 $25~\mu L$ of freshly prepared Charcoal waste stream and 5 μL of 10% HD, HN and L solutions

Key for HGPs #379, 380, 387, and 388 dosed 8/29/1996. Exposure duration - 1 hr.

Animal # 379

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Charcoal waste stream
Е	5 μL of 10% HN in CHCl ₃
F	25 μL of Charcoal waste stream

Site	Treatment
A	5 μL of 10% HN in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% L in CHCl ₃
D	25 μL of Charcoal waste stream
Е	5 μL of 10% HD in CHCl ₃
F	25 μL of Charcoal waste stream

Site	Treatment
A	5 μL of 10% HD in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% HN in CHCl ₃
D	25 μL of Charcoal waste stream
E	5 μL of 10% L in CHCl ₃
F	25 μL of Charcoal waste stream

Site	Treatment
A	5 μL of 10% L in CHCl ₃
В	25 μL of Charcoal waste stream
С	5 μL of 10% HD in CHCl ₃
D	25 μL of Charcoal waste stream
Е	5 μL of 10% HN in CHCl ₃
F	25 μL of Charcoal waste stream

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MREF Task 95-38 G1555-38A

Animal # 379 Site		A	В	С	D	E	F	
Histopathology Markers:								
Microblister	4	0	2	0	3	0		
Epidermal Necrosis	4	1**	4	1**	4	1**		
Follicular Necrosis	4	1**	4	1**	4	1**		
Dermalal Necrosis	0	0	0*	0	0	0		
Vascular Necrosis	0	0	0	0	0	0		
Hemorrhage	2	0	2	0	1	0		
Pustular Epidermitis	0	0	0	0	0	0		
Notes: *moderate derma edema; **random single necrosis noted	mod dermal inflam	min dermal inflam	mild dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam		
Animal # 380	Site	A	В	C	D	E	F	
Histopathology Markers	:						·	
Microblister		4	0	4	0	3	0	
Epidermal Necrosis		4	1*	4	71*	4	0	
Follicular Necrosis		4	1*	4	1*	4	1*	
Dermal: Necrosis		1	.0	2**	0	3**	0	
Vascular Necrosis	0	0	0	0	0	0		
Hemorrhage		0	0	1	0	0	0	
Pustular Epidermitis	0	0	0	0	0	0		
Notes: *random single cell necrosis **mod dermal edema; focal ulcer in area of necrosis		mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	mod dermal inflam	min dermal inflam	

Note: Some normal (unaffected) skin present at one or both ends of each section where lesions were present.

Histopathological Markers: Degree of Severity Grading Scale

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0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

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Animal # 387 Site		A	В	С	D	E	F
Histopathology Marker	s:						
Microblister	3	0	2	0	4	0	
Epidermal. Necrosis	4	1**	4	1**	4	1**	
Follicular Necrosis	4	1**	3	1**	4	1**	
Dermal Necrosis	0	0	0	0	0*	0	
Vascular Necrosis	0	0	0	0	0	0	
Hemorrhage	0	0	0	0	2	0	
Pustular Epidermitis	0	0	0	0	0	1	
Notes: *moderate dermal edema; **random single-cell necrosis noted		mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam	mod dermal inflam
Animal # 388	Α	В	С	D	Е	F	
Histopathology Markers	s:						·
Histopathology Markers Microblister	s:	4	0	4	0	3	0
	S:	4	0	4	0	3 4	0
Microblister	s:						
Microblister Epidermal Necrosis	s:	4	1*	4	1*	4	1*
Microblister Epidermal Necrosis Follicular Necrosis	S:	4	1*	4	1*	4 2	1*
Microblister Epidermal Necrosis Follicular Necrosis Dermal Necrosis	S:	4 4 0**	1* 1* 0	4 4 0**	1* 1* 0	4 2 0	1* 1* 0
Microblister Epidermal Necrosis Follicular Necrosis Dermal Necrosis Vascular Necrosis	S:	4 4 0** 0	1* 1* 0 0	4 4 0**	1* . 1* . 0	4 2 0 0	1* 1* 0 0

Histopathological Markers: Degree of Severity Grading Scale

9/9/96

0 = Normal; 1 = Minimal; 2 = Intermediate; 3 = Moderate; 4 = Severe

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